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(54) Title: NUCLEIC ACID SEQUENCE ENCODING BETA-C-4-OXYGENASE FROM HAEMATOCOCCUS PLUVIALIS FOR THE BIOSYNTHESIS OF ASTAXANTHIN

(57) Abstract

The present invention relates, in general, to a biotechnological method for production of (3S,3'S) astaxanthin. In particular, the present invention relates to a peptide having a β -C-4-oxygenase activity; a DNA segment coding for this peptide; an RNA segment coding for this peptide; a recombinant DNA molecule comprising a vector and the DNA segment; a host cell or organism containing the above described recombinant DNA molecule or DNA segment; and to a method of biotechnologically producing (3S,3'S) astaxanthin or a food additive containing (3S,3'S) astaxanthin, using the host.

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NUCLEIC ACID SEQUENCE ENCODING BETA-C-4-OXYGENASE FROM HAEMATOCOCCUS PLUVIALIS FOR THE BIOSYNTHESIS OF ASTAXANTHIN

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FIELD AND BACKGROUND OF THE INVENTION

The present invention relates, in general, to a biotechnological method for production of (3S,3'S) astaxanthin. In particular, the present invention relates to a peptide having a β -C-4-oxygenase activity; a DNA segment coding for this peptide; an RNA segments coding for this peptide; a recombinant DNA molecule comprising a vector and the DNA segment; a host cell or organism containing the above described recombinant DNA molecule or DNA segment; and to a method of biotechnologically producing (3S,3'S) astaxanthin or a food additive containing (3S,3'S) astaxanthin, using the host.

Carotenoids, such as astaxanthin, are natural pigments that are responsible for many of the yellow, orange and red colors seen in living organisms. Carotenoids are widely distributed in nature and have, in various living systems, two main biological functions: they serve as light-harvesting pigments in photosynthesis, and they protect against photooxidative damage. These and additional biological functions of carotenoids, their important industrial role, and their biosynthesis are discussed hereinbelow.

As part of the light-harvesting antenna, carotenoids can absorb photons and transfer the energy to chlorophyll, thus assisting in the harvesting of light in the range of 450 - 570 nm [see, Cogdell RJ and Frank HA (1987) How carotenoids function in photosynthestic bacteria. Biochim Biophys Acta 895: 63-79; Cogdell R (1988) The function of pigments in chloroplasts. In: Goodwin TW (ed) Plant Pigments, pp 183-255. Academic Press, London; Frank HA, Violette CA, Trautman JK, Shreve AP, Owens TG and Albrecht AC (1991) Carotenoids in photosynthesis: structure and photochemistry. Pure Appl Chem 63: 109-114; Frank HA, Farhoosh R, Decoster B and Christensen RL (1992) Molecular features that control the efficiency of carotenoid-to-chlorophyll energy photosynthesis. In: Murata N (ed) Research in Photosynthesis, Vol I, pp 125-128. Kluwer, Dordrecht; and, Cogdell RJ and Gardiner AT (1993) Functions of carotenoids in photosynthesis. Meth Enzymol 214: 185-193]. carotenoids are integral constituents of the protein-pigment complexes of the lightharvesting antennae in photosynthetic organisms, they are also important components of the photosynthetic reaction centers.

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Most of the total carotenoids is located in the light harvesting complex II [Bassi R, Pineaw B, Dainese P and Marquartt J (1993) Carotenoid binding proteins Eur J Biochem 212: 297-302]. The identities of the of photosystem II. photosynthetically active carotenoproteins and their precise location in lightharvesting systems are not known. Carotenoids in photochemically active chlorophyll-protein complexes of the thermophilic cyanobacterium Synechococcus sp. were investigated by linear dichroism spectroscopy of oriented samples [see, Breton J and Kato S (1987) Orientation of the pigments in photosystem II: lowtemperature linear-dichroism study of a core particle and of its chlorophyll-protein subunits isolated from Synechococcus sp. Biochim Biophys Acta 892: 99-107]. These complexes contained mainly a \beta-carotene pool absorbing around 505 and 470 nm, which is oriented close to the membrane plane. In photochemically inactive chlorophyll-protein complexes, the \beta-carotene absorbs around 495 and 465 nm, and the molecules are oriented perpendicular to the membrane plane.

Evidence that carotenoids are associated with cyanobacterial photosystem (PS) II has been described [see, Suzuki R and Fujita Y (1977) Carotenoid photobleaching induced by the action of photosynthetic reaction center II: DCMU sensitivity. Plant Cell Physiol 18: 625-631; and, Newman PJ and Sherman LA (1978) Isolation and characterization of photosystem I and II membrane particles from the blue-green alga Synechococcus cedrorum. Biochim Biophys Acta 503: 343-361]. There are two β -carotene molecules in the reaction center core of PS II [see, Ohno T, Satoh K and Katoh S (1986) Chemical composition of purified oxygen-evolving complexes from the thermophilic cyanobacterium Synechococcus sp. Biochim Biophys Acta 852: 1-8; Gounaris K, Chapman DJ and Barber J (1989) Isolation and characterization of a D1/D2/cytochrome b-559 complex from Synechocystis PCC6803. Biochim Biophys Acta 973: 296-301; and, Newell RW, van Amerongen H, Barber J and van Grondelle R (1993) Spectroscopic characterization of the reaction center of photosystem II using polarized light: Evidence for β-carotene excitors in PS II reaction centers. Biochim Biophys Acta 1057: 232-238] whose exact function(s) is still obscure [reviewed by Satoh K (1992) Structure and function of PS II reaction center. In: Murata N (ed) Research in Photosynthesis, Vol. II, pp. 3-12. Kluwer, Dordrecht]. It was demonstrated that these two coupled \(\beta\)-carotene molecules protect chlorophyll P680 from photodamage in isolated PS II reaction centers [see, De Las Rivas J, Telfer A and Barber J (1993) 2-coupled β-carotene molecules protect P680 from photodamage in isolated PS II reaction centers. Biochim. Biophys. Acta 1142: 155-164], and this may be related to the protection against degradation of the D1 subunit of PS II [see, Sandmann G (1993) Genes and enzymes involved in the desaturation

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reactions from phytoene to lycopene. (abstract), 10th International Symposium on Carotenoids, Trondheim CL1-2]. The light-harvesting pigments of a highly purified, oxygen-evolving PS II complex of the thermophilic cyanobacterium *Synechococcus* sp. consists of 50 chlorophyll *a* and 7 β-carotene, but no xanthophyll, molecules [see, Ohno T, Satoh K and Katoh S (1986) Chemical composition of purified oxygen-evolving complexes from the thermophilic cyanobacterium *Synechococcus* sp. Biochim Biophys Acta 852: 1-8]. β-carotene was shown to play a role in the assembly of an active PS II in green algae [see, Humbeck K, Romer S and Senger H (1989) Evidence for the essential role of carotenoids in the assembly of an active PS II. Planta 179: 242-250].

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Isolated complexes of PS I from Phormidium luridum, which contained 40 chlorophylls per P700, contained an average of 1.3 molecules of β-carotene [see, Thornber JP, Alberte RS, Hunter FA, Shiozawa JA and Kan KS (1976) The organization of chlorophyll in the plant photosynthetic unit. Brookhaven Symp Biology 28: 132-148]. In a preparation of PS I particles from Synechococcus sp. strain PCC 6301, which contained 130 ± 5 molecules of antenna chlorophylls per P700, 16 molecules of carotenoids were detected [see, Lundell DJ, Glazer AN, Melis A and Malkin R (1985) Characterization of a cyanobacterial photosystem I complex. J Biol Chem 260: 646-654]. A substantial content of β -carotene and the xanthophylls cryptoxanthin and isocryptoxanthin were detected in PS I pigmentprotein complexes of the thermophilic cyanobacterium Synechococcus elongatus Isee, Coufal J, Hladik J and Sofrova D (1989) The carotenoid content of photosystem 1 pigment-protein complexes of the cyanobacterium Synechococcus elongatus. Photosynthetica 23: 603-616]. A subunit protein-complex structure of PS I from the thermophilic cyanobacterium Synechococcus sp., which consisted of four polypeptides (of 62, 60, 14 and 10 kDa), contained approximately 10 βcarotene molecules per P700 [see, Takahashi Y, Hirota K and Katoh S (1985) Multiple forms of P700-chlorophyll a-protein complexes from Synechococcus sp.: the iron, quinone and carotenoid contents. Photosynth Res 6: 183-192]. carotenoid is exclusively bound to the large polypeptides which carry the functional and antenna chlorophyll a. The fluorescence excitation spectrum of these complexes suggested that β-carotene serves as an efficient antenna for PS I.

As mentioned, an additional essential function of carotenoids is to protect against photooxidation processes in the photosynthetic apparatus that are caused by the excited triplet state of chlorophyll. Carotenoid molecules with π -electron conjugation of nine or more carbon-carbon double bonds can absorb triplet-state energy from chlorophyll and thus prevent the formation of harmful singlet-state oxygen radicals. In *Synechococcus* sp. the triplet state of carotenoids was

monitored in closed PS II centers and its rise kinetics of approximately 25 nanoseconds is attributed to energy transfer from chlorophyll triplets in the antenna [see, Schlodder E and Brettel K (1988) Primary charge separation in closed photosystem II with a lifetime of 11 nanoseconds. Flash-absorption spectroscopy with oxygen-evolving photosystem II complexes from *Synechococcus*. Biochim Biophys Acta 933: 22-34]. It is conceivable that this process, that has a lower yield compared to the yield of radical-pair formation, plays a role in protecting chlorophyll from damage due to over-excitation.

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The protective role of carotenoids *in vivo* has been elucidated through the use of bleaching herbicides such as norflurazon that inhibit carotenoid biosynthesis in all organisms performing oxygenic photosynthesis [reviewed by Sandmann G and Boger P (1989) Inhibition of carotenoid biosynthesis by herbicides. In: Boger P and Sandmann G (Eds.) Target Sites of Herbicide Action, pp 25-44. CRC Press, Boca Raton, Florida]. Treatment with norflurazon in the light results in a decrease of both carotenoid and chlorophyll levels, while in the dark, chlorophyll levels are unaffected. Inhibition of photosynthetic efficiency in cells of *Oscillatoria agardhii* that were treated with the pyridinone herbicide, fluridone, was attributed to a decrease in the relative abundance of myxoxanthophyll, zeaxanthin and β -carotene, which in turn caused photooxidation of chlorophyll molecules [see, Canto de Loura I, Dubacq JP and Thomas JC (1987) The effects of nitrogen deficiency on pigments and lipids of cianobacteria. Plant Physiol 83: 838-843].

It has been demonstrated in plants that zeaxanthin is required to dissipate, in a nonradiative manner, the excess excitation energy of the antenna chlorophyll [see, Demmig-Adams B (1990) Carotenoids and photoprotection in plants: a role for the xanthophyll zeaxanthin. Biochim Biophys Acta 1020: 1-24; and, Demmig-Adams B and Adams WW III (1990) The carotenoid zeaxanthin and high-energystate quenching of chlorophyll fluorescence. Photosynth Res 25: 187-197]. In algae and plants a light-induced deepoxidation of violaxanthin to yield zeaxanthin, is related to photoprotection processes [reviewed by Demmig-Adams B and Adams WW III (1992) Photoprotection and other responses of plants to high light stress. Ann Rev Plant Physiol Plant Mol Biol 43: 599-626]. The light-induced deepoxidation of violaxanthin and the reverse reaction that takes place in the dark, are known as the "xanthophyll cycle" [see, Demmig-Adams B and Adams WW III (1992) Photoprotection and other responses of plants to high light stress. Ann Rev Plant Physiol Plant Mol Biol 43: 599-626]. Cyanobacterial lichens, that do not contain any zeaxanthin and that probably are incapable of radiationless energy dissipation, are sensitive to high light intensity; algal lichens that contain zeaxanthin are more resistant to high-light stress [see, Demmig-Adams B, Adams

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WW III, Green TGA, Czygan FC and Lange OL (1990) Differences in the susceptibility to light stress in two lichens forming a phycosymbiodeme, one partner possessing and one lacking the xanthophyll cycle. Oecologia 84: 451-456: Demmig-Adams B and Adams WW III (1993) The xanthophyll cycle, protein turnover, and the high light tolerance of sun-acclimated leaves. Plant Physiol 103: 1413-1420; and, Demmig-Adams B (1990) Carotenoids and photoprotection in plants: a role for the xanthophyll zeaxanthin. Biochim Biophys Acta 1020: 1-24]. In contrast to algae and plants, cyanobacteria do not have a xanthophyll cycle. However, they do contain ample quantities of zeaxanthin and other xanthophylls that can support photoprotection of chlorophyll.

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Several other functions have been ascribed to carotenoids. The possibility that carotenoids protect against damaging species generated by near ultra-violet (UV) irradiation is suggested by results describing the accumulation of β-carotene in a UV-resistant mutant of the cyanobacterium Gloeocapsa alpicola [see, Buckley CE and Houghton JA (1976) A study of the effects of near UV radiation on the pigmentation of the blue-green alga Gloeocapsa alpicola. Arch Microbiol 107: 93-97]. This has been demonstrated more elegantly in Escherichia coli cells that produce carotenoids [see, Tuveson RW and Sandmann G (1993) Protection by cloned carotenoid genes expressed in Escherichia coli against phototoxic molecules activated by near-ultraviolet light. Meth Enzymol 214: 323-330]. Due to their ability to quench oxygen radical species, carotenoids are efficient antioxidants and thereby protect cells from oxidative damage. This function of carotenoids is important in virtually all organisms [see, Krinsky NI (1989) Antioxidant functions of carotenoids. Free Radical Biol Med 7: 617-635; and, Palozza P and Krinsky NI (1992) Antioxidant effects of carotenoids in vivo and in vitro - an overview. Meth Enzymol 213: 403-420]. Other cellular functions could be affected by carotenoids, even if indirectly. Although carotenoids in cyanobacteria are not the major photoreceptors for phototaxis, an influence of carotenoids on phototactic reactions, that have been observed in Anabaena variabilis, was attributed to the removal of singlet oxygen radicals that may act as signal intermediates in this system [see, Nultsch W and Schuchart H (1985) A model of the phototactic reaction chain of cyanobacterium Anabaena variabilis. Arch Microbiol 142: 180-184].

In flowers and fruits carotenoids facilitate the attraction of pollinators and dispersal of seeds. This latter aspect is strongly associated with agriculture. The type and degree of pigmentation in fruits and flowers are among the most important traits of many crops. This is mainly since the colors of these products

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often determine their appeal to the consumers and thus can increase their market worth.

Carotenoids have important commercial uses as coloring agents in the food industry since they are non-toxic [see, Bauernfeind JC (1981) Carotenoids as colorants and vitamin A precursors. Academic Press, London]. The red color of the tomato fruit is provided by lycopene which accumulates during fruit ripening in chromoplasts. Tomato extracts, which contain high content (over 80% dry weight) of lycopene, are commercially produced worldwide for industrial use as food colorant. Furthermore, the flesh, feathers or eggs of fish and birds assume the color of the dietary carotenoid provided, and thus carotenoids are frequently used in dietary additives for poultry and in aquaculture. Certain cyanobacterial species, for example *Spirulina* sp. [see, Sommer TR, Potts WT and Morrissy NM (1990) Recent progress in processed microalgae in aquaculture. Hydrobiologia 204/205: 435-443], are cultivated in aquaculture for the production of animal and human food supplements. Consequently, the content of carotenoids, primarily of β-carotene, in these cyanobacteria has a major commercial implication in biotechnology.

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Most carotenoids are composed of a C40 hydrocarbon backbone, constructed from eight C5 isoprenoid units and contain a series of conjugated double bonds. Carotenes do not contain oxygen atoms and are either linear or cyclized molecules containing one or two end rings. Xanthophylls are oxygenated derivatives of carotenes. Various glycosilated carotenoids and carotenoid esters have been identified. The C40 backbone can be further extended to give C45 or C50 carotenoids, or shortened yielding apocarotenoids. Some nonphotosynthetic bacteria also synthesize C30 carotenoids. General background on carotenoids can be found in Goodwin TW (1980) The Biochemistry of the Carotenoids, Vol. 1, 2nd Ed. Chapman and Hall, New York; and in Goodwin TW and Britton G (1988) Distribution and analysis of carotenoids. In: Goodwin TW (ed) Plant Pigments, pp 62-132. Academic Press, New York.

More than 640 different naturally-occurring carotenoids have been so far characterized, hence, carotenoids are responsible for most of the various shades of yellow, orange and red found in microorganisms, fungi, algae, plants and animals. Carotenoids are synthesized by all photosynthetic organisms as well as several nonphotosynthetic bacteria and fungi, however they are also widely distributed through feeding throughout the animal kingdom.

Carotenoids are synthesized de novo from isoprenoid precursors only in photosynthetic organisms and some microorganisms, they typically accumulate in

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protein complexes in the photosynthetic membrane, in the cell membrane and in the cell wall.

As detailed in Figure 1, in the biosynthesis pathway of β-carotene, four enzymes convert geranylgeranyl pyrophosphate of the central isoprenoid pathway to β-carotene. Carotenoids are produced from the general isoprenoid biosynthetic pathway. While this pathway has been known for several decades, only recently, and mainly through the use of genetics and molecular biology, have some of the molecular mechanisms involved in carotenoids biogenesis, been elucidated. This is due to the fact that most of the enzymes which take part in the conversion of phytoene to carotenes and xanthophylls are labile, membrane-associated proteins that lose activity upon solubilization [see, Beyer P, Weiss G and Kleinig H (1985) Solubilization and reconstitution of the membrane-bound carotenogenic enzymes from daffodile chromoplasts. Eur J Biochem 153: 341-346; and, Bramley PM (1985) The in vitro biosynthesis of carotenoids. Adv Lipid Res 21: 243-279]. However, solubilization of carotenogenic enzymes from Synechocystis sp. strain PCC 6714 that retain partial activity has been reported [see, Bramley PM and Sandmann G (1987) Solubilization of carotenogenic enzyme of Aphanocapsa. Phytochem 26: 1935-1939]. There is no genuine in vitro system for carotenoid biosynthesis which enables a direct essay of enzymatic activities. A cell-free carotenogenic system has been developed [see, Clarke IE, Sandmann G, Bramley PM and Boger P (1982) Carotene biosynthesis with isolated photosynthetic membranes. FEBS Lett 140: 203-206] and adapted for cyanobacteria [see, Sandmann G and Bramley PM (1985) Carotenoid biosynthesis by Aphanocapsa homogenates coupled to a phytoene-generating system from Phycomyces blakesleeanus. Planta 164: 259-263; and, Bramley PM and Sandmann G (1985) In and in vivo biosynthesis of xanthophylls by the cyanobacterium Aphanocapsa. Phytochem 24: 2919-2922]. Reconstitution of phytoene desaturase from Synechococcus sp. strain PCC 7942 in liposomes was achieved following purification of the polypeptide, that had been expressed in Escherichia coli [see, Fraser PD, Linden H and Sandmann G (1993) Purification and reactivation of recombinant Synechococcus phytoene desaturase from an overexpressing strain of Escherichia coli. Biochem J 291: 687-692].

Referring now to Figure 1, carotenoids are synthesized from isoprenoid precursors. The central pathway of isoprenoid biosynthesis may be viewed as beginning with the conversion of acetyl-CoA to mevalonic acid. D³-isopentenyl pyrophosphate (IPP), a C₅ molecule, is formed from mevalonate and is the building block for all long-chain isoprenoids. Following isomerization of IPP to dimethylallyl pyrophosphate (DMAPP), three additional molecules of IPP are

combined to yield the C₂₀ molecule, geranylgeranyl pyrophosphate (GGPP). These 1'-4 condensation reactions are catalyzed by prenyl transferases [see, Kleinig H (1989) The role of plastids in isoprenoid biosynthesis. Ann Rev Plant Physiol Plant Mol Biol 40: 39-59]. There is evidence in plants that the same enzyme, GGPP synthase, carries out all the reactions from DMAPP to GGPP [see, Dogbo O and Camara B (1987) Purification of isopentenyl pyrophosphate isomerase and geranylgeranyl pyrophosphate synthase from *Capsicum* chromoplasts by affinity chromatography. Biochim Biophys Acta 920: 140-148; and, Laferriere A and Beyer P (1991) Purification of geranylgeranyl diphosphate synthase from *Sinapis alba* etioplasts. Biochim Biophys Acta 216: 156-163].

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The first step that is specific for carotenoid biosynthesis is the head-to-head condensation of two molecules of GGPP to produce prephytoene pyrophosphate (PPPP). Following removal of the pyrophosphate, GGPP is converted to 15-cisphytoene, a colorless C40 hydrocarbon molecule. This two-step reaction is catalyzed by the soluble enzyme, phytoene synthase, an enzyme encoded by a single gene (crtB), in both cyanobacteria and plants [see, Chamovitz D, Misawa N, Sandmann G and Hirschberg J (1992) Molecular cloning and expression in Escherichia coli of a cyanobacterial gene coding for phytoene synthase, a carotenoid biosynthesis enzyme. FEBS Lett 296: 305-310; Ray JA, Bird CR, Maunders M, Grierson D and Schuch W (1987) Sequence of pTOM5, a ripening related cDNA from tomato. Nucl Acids Res 15: 10587-10588; Camara B (1993) Plant phytoene synthase complex - component 3 enzymes, immunology, and biogenesis. Meth Enzymol 214: 352-365]. All the subsequent steps in the pathway Four desaturation (dehydrogenation) reactions convert occur in membranes. phytoene to lycopene via phytofluene, ζ-carotene, and neurosporene. desaturation increases the number of conjugated double bonds by two such that the number of conjugated double bonds increases from three in phytoene to eleven in lycopene.

Relatively little is known about the molecular mechanism of the enzymatic dehydrogenation of phytoene [see, Jones BL and Porter JW (1986) Biosynthesis of carotenes in higher plants. CRC Crit Rev Plant Sci 3: 295-324; and, Beyer P, Mayer M and Kleinig H (1989) Molecular oxygen and the state of geometric iosomerism of intermediates are essential in the carotene desaturation and cyclization reactions in daffodil chromoplasts. Eur J Biochem 184: 141-150]. It has been established that in cyanobacteria, algae and plants the first two desaturations, from 15-cis-phytoene to ζ -carotene, are catalyzed by a single membrane-bound enzyme, phytoene desaturase [see, Jones BL and Porter JW (1986) Biosynthesis of carotenes in higher plants. CRC Crit Rev Plant Sci 3: 295-

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324; and, Beyer P, Mayer M and Kleinig H (1989) Molecular oxygen and the state of geometric iosomerism of intermediates are essential in the carotene desaturation and cyclization reactions in daffodil chromoplasts. Eur J Biochem 184: 141-150]. Since the ζ-carotene product is mostly in the all-trans configuration, a cis-trans isomerization is presumed at this desaturation step. The primary structure of the phytoene desaturase polypeptide in cyanobacteria is conserved (over 65% identical residues) with that of algae and plants [see, Pecker I, Chamovitz D, Linden H, Sandmann G and Hirschberg J (1992) A single polypeptide catalyzing the conversion of phytoene to ζ-carotene is transcriptionally regulated during tomato fruit ripening. Proc Natl Acad Sci USA 89: 4962-4966; Pecker I, Chamovitz D. Mann V, Sandmann G, Boger P and Hirschberg J (1993) Molecular characterization of carotenoid biosynthesis in plants: the phytoene desaturase gene in tomato. In: Murata N (ed) Research in Photosynthesis, Vol III, pp 11-18. Kluwer, Dordrectht]. Moreover, the same inhibitors block phytoene desaturase in the two systems [see, Sandmann G and Boger P (1989) Inhibition of carotenoid biosynthesis by herbicides. In: Boger P and Sandmann G (eds) Target Sites of Herbicide Action, pp 25-44. CRC Press, Boca Raton, Florida]. Consequently, it is very likely that the enzymes catalyzing the desaturation of phytoene and phytofluene in cyanobacteria and plants have similar biochemical and molecular properties, that are distinct from those of phytoene desaturases in other microorganisms. One such a difference is that phytoene desaturases from Rhodobacter capsulatus, Erwinia sp. or fungi convert phytoene to neurosporene, lycopene, or 3,4-dehydrolycopene, respectively.

Desaturation of phytoene in daffodil chromoplasts [see, Beyer P, Mayer M and Kleinig H (1989) Molecular oxygen and the state of geometric iosomerism of intermediates are essential in the carotene desaturation and cyclization reactions in daffodil chromoplasts. Eur J Biochem 184: 141-150], as well as in a cell free system of Synechococcus sp. strain PCC 7942 [see, Sandmann G and Kowalczyk S (1989) In vitro carotenogenesis and characterization of the phytoene desaturase reaction in Anacystis. Biochem Biophys Res Com 163: 916-921], is dependent on molecular oxygen as a possible final electron acceptor, although oxygen is not directly involved in this reaction. A mechanism of dehydrogenase-electron transferase was supported in cyanobacteria over dehydrogenation mechanism of dehydrogenase-monooxygenase [see, Sandmann G and Kowalczyk S (1989) In vitro carotenogenesis and characterization of the phytoene desaturase reaction in Anacystis. Biochem Biophys Res Com 163: 916-921]. A conserved FAD-binding motif exists in all phytoene desaturases whose primary structures have been analyzed [see, Pecker I, Chamovitz D, Linden H, Sandmann G and Hirschberg J

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(1992) A single polypeptide catalyzing the conversion of phytoene to ζ -carotene is transcriptionally regulated during tomato fruit ripening. Proc Natl Acad Sci USA 89: 4962-4966; Pecker I, Chamovitz D, Mann V, Sandmann G, Boger P and Hirschberg J (1993) Molecular characterization of carotenoid biosynthesis in plants: the phytoene desaturase gene in tomato. In: Murata N (ed) Research in Photosynthesis, Vol III, pp 11-18. Kluwer, Dordrectht]. The phytoene desaturase enzyme in pepper was shown to contain a protein-bound FAD [see, Hugueney P, Romer S, Kuntz M and Camara B (1992) Characterization and molecular cloning of a flavoprotein catalyzing the synthesis of phytofluene and ζ -carotene in Capsicum chromoplasts. Eur J Biochem 209: 399-407]. Since phytoene desaturase is located in the membrane, an additional, soluble redox component is predicted. This hypothetical component could employ NAD(P)+, as suggested [see, Mayer MP, Nievelstein V and Beyer P (1992) Purification and characterization of a chromoplasts from oxidoreductase dependent NADPH pseudonarcissus - a redox-mediator possibly involved in carotene desaturation. Plant Physiol Biochem 30: 389-398] or another electron and hydrogen carrier, such as a quinone. The cellular location of phytoene desaturase in Synechocystis sp. strain PCC 6714 and Anabaena variabilis strain ATCC 29413 was determined with specific antibodies to be mainly (85%) in the photosynthetic thylakoid membranes [see, Serrano A, Gimenez P, Schmidt A and Sandmann G (1990) Immunocytochemical localization and functional determination of phytoene desaturase in photoautotrophic prokaryotes. J Gen Microbiol 136: 2465-2469].

In cyanobacteria algae and plants ζ -carotene is converted to lycopene via neurosporene. Very little is known about the enzymatic mechanism, which is predicted to be carried out by a single enzyme [see, Linden H, Vioque A and Sandmann G (1993) Isolation of a carotenoid biosynthesis gene coding for ζ -carotene desaturase from *Anabaena* PCC 7120 by heterologous complementation. FEMS Microbiol Lett 106: 99-104]. The deduced amino acid sequence of ζ -carotene desaturase in *Anabaena* sp. strain PCC 7120 contains a dinucleotide-binding motif that is similar to the one found in phytoene desaturase.

Two cyclization reactions convert lycopene to β-carotene. Evidence has been obtained that in *Synechococcus* sp. strain PCC 7942 [see, Cunningham FX Jr, Chamovitz D, Misawa N, Gantt E and Hirschberg J (1993) Cloning and functional expression in *Escherichia coli* of a cyanobacterial gene for lycopene cyclase, the enzyme that catalyzes the biosynthesis of β-carotene. FEBS Lett 328: 130-138], as well as in plants [see, Camara B and Dogbo O (1986) Demonstration and solubilization of lycopene cyclase from *Capsicum* chromoplast membranes. Plant Physiol 80: 172-184], these two cyclizations are catalyzed by a single enzyme.

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lycopene cyclase. This membrane-bound enzyme is inhibited by the triethylamine compounds, CPTA and MPTA [see, Sandmann G and Boger P (1989) Inhibition of carotenoid biosynthesis by herbicides. In: Boger P and Sandmann G (eds) Target Sites of Herbicide Action, pp 25-44. CRC Press, Boca Raton, Floridal. Cyanobacteria carry out only the β-cyclization and therefore do not contain εcarotene, δ -carotene and α -carotene and their oxygenated derivatives. The β -ring is formed through the formation of a "carbonium ion" intermediate when the C-1,2 double bond at the end of the linear lycopene molecule is folded into the position of the C-5,6 double bond, followed by a loss of a proton from C-6. No cyclic carotene has been reported in which the 7,8 bond is not a double bond. Therefore, full desaturation as in lycopene, or desaturation of at least half-molecule as in neurosporene, is essential for the reaction. Cyclization of lycopene involves a dehydrogenation reaction that does not require oxygen. The cofactor for this reaction is unknown. A dinucleotide-binding domain was found in the lycopene cyclase polypeptide of Synechococcus sp. strain PCC 7942, implicating NAD(P) or FAD as coenzymes with lycopene cyclase.

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The addition of various oxygen-containing side groups, such as hydroxy-, methoxy-, oxo-, epoxy-, aldehyde or carboxylic acid moieties, form the various xanthophyll species. Little is known about the formation of xanthophylls. Hydroxylation of β -carotene requires molecular oxygen in a mixed-function oxidase reaction.

Clusters of genes encoding the enzymes for the entire pathway have been cloned from the purple photosynthetic bacterium Rhodobacter capsulatus [see, Armstrong GA, Alberti M, Leach F and Hearst JE (1989) Nucleotide sequence, organization, and nature of the protein products of the carotenoid biosynthesis gene cluster of Rhodobacter capsulatus. Mol Gen Genet 216: 254-268] and from the nonphotosynthetic bacteria Erwinia herbicola [see, Sandmann G, Woods WS and Tuveson RW (1990) Identification of carotenoids in Erwinia herbicola and in transformed Escherichia coli strain. FEMS Microbiol Lett 71: 77-82; Hundle BS, Beyer P, Kleinig H, Englert H and Hearst JE (1991) Carotenoids of Erwinia herbicola and an Escherichia coli HB101 strain carrying the Erwinia herbicola carotenoid gene cluster. Photochem Photobiol 54: 89-93; and, Schnurr G, Schmidt A and Sandmann G (1991) Mapping of a carotenogenic gene cluster from Erwinia herbicola and functional identification of six genes. FEMS Microbiol Lett 78: 157-162] and Erwinia uredovora [see, Misawa N, Nakagawa M, Kobayashi K, Yamano S, Izawa I, Nakamura K and Harashima K (1990) Elucidation of the Erwinia uredovora carotenoid biosynthetic pathway by functional analysis of gene products in Escherichia coli. J Bacteriol 172: 6704-6712]. Two genes, al-3

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for GGPP synthase [see, Nelson MA, Morelli G, Carattoli A, Romano N and Macino G (1989) Molecular cloning of a *Neurospora crassa* carotenoid biosynthetic gene (*albino-3*) regulated by blue light and the products of the white collar genes. Mol Cell Biol 9: 1271-1276; and, Carattoli A, Romano N, Ballario P, Morelli G and Macino G (1991) The *Neurospora crassa* carotenoid biosynthetic gene (albino 3). J Biol Chem 266: 5854-5859] and *al-1* for phytoene desaturase [see, Schmidhauser TJ, Lauter FR, Russo VEA and Yanofsky C (1990) Cloning sequencing and photoregulation of *al-1*, a carotenoid biosynthetic gene of *Neurospora crassa*. Mol Cell Biol 10: 5064-5070] have been cloned from the fungus *Neurospora crassa*. However, attempts at using these genes as heterologous molecular probes to clone the corresponding genes from cyanobacteria or plants were unsuccessful due to lack of sufficient sequence similarity.

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The first "plant-type" genes for carotenoid synthesis enzyme were cloned from cyanobacteria using a molecular-genetics approach. In the first step towards cloning the gene for phytoene desaturase, a number of mutants that are resistant to the phytoene-desaturase-specific inhibitor, norflurazon, were isolated in Synechococcus sp. strain PCC 7942 [see, Linden H, Sandmann G, Chamovitz D, Hirschberg J and Boger P (1990) Biochemical characterization of Synechococcus mutants selected against the bleaching herbicide norflurazon. Pestic Biochem Physiol 36: 46-51]. The gene conferring norflurazon-resistance was then cloned by transforming the wild-type strain to herbicide resistance [see, Chamovitz D, Pecker I and Hirschberg J (1991) The molecular basis of resistance to the herbicide norflurazon. Plant Mol Biol 16: 967-974; Chamovitz D, Pecker I, Sandmann G, Boger P and Hirschberg J (1990) Cloning a gene for norflurazon resistance in cyanobacteria. Z Naturforsch 45c: 482-486]. Several lines of evidence indicated that the cloned gene, formerly called pds and now named crtP, codes for phytoene desaturase. The most definitive one was the functional expression of phytoene desaturase activity in transformed Escherichia coli cells [see, Linden H, Misawa N, Chamovitz D, Pecker I, Hirschberg J and Sandmann G (1991) Functional complementation in Escherichia coli of different phytoene desaturase genes and analysis of accumulated carotenes. Z Naturforsch 46c: 1045-1051; and. Pecker I, Chamovitz D, Linden H, Sandmann G and Hirschberg J (1992) A single catalyzing the conversion of phytoene to ζ -carotene is polypeptide transcriptionally regulated during tomato fruit ripening. Proc Natl Acad Sci USA 89: 4962-4966]. The crtP gene was also cloned from Synechocystis sp. strain PCC 6803 by similar methods [see, Martinez-Ferez IM and Vioque A (1992) Nucleotide sequence of the phytoene desaturase gene from Synechocystis sp. PCC

6803 and characterization of a new mutation which confers resistance to the herbicide norflurazon. Plant Mol Biol 18: 981-983].

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The cyanobacterial crtP gene was subsequently used as a molecular probe for cloning the homologous gene from an alga [see, Pecker I, Chamovitz D, Mann V, Sandmann G, Boger P and Hirschberg J (1993) Molecular characterization of carotenoid biosynthesis in plants: the phytoene desaturase gene in tomato. In: Murata N (ed) Research in Photosynthesis, Vol III, pp 11-18. Kluwer, Dordrectht] and higher plants [see, Bartley GE, Viitanen PV, Pecker I, Chamovitz D, Hirschberg J and Scolnik PA (1991) Molecular cloning and expression in photosynthetic bacteria of a soybean cDNA coding for phytoene desaturase, an enzyme of the carotenoid biosynthesis pathway. Proc Natl Acad Sci USA 88: 6532-6536; and, Pecker I, Chamovitz D, Linden H, Sandmann G and Hirschberg J (1992) A single polypeptide catalyzing the conversion of phytoene to ζ -carotene is transcriptionally regulated during tomato fruit ripening. Proc Natl Acad Sci USA 89: 4962-4966]. The phytoene desaturases in Synechococcus sp. strain PCC 7942 and Synechocystis sp. strain PCC 6803 consist of 474 and 467 amino acid residues, respectively, whose sequences are highly conserved (74% identities and 86% similarities). The calculated molecular mass is 51 kDa and, although it is slightly hydrophobic (hydropathy index -0.2), it does not include a hydrophobic region which is long enough to span a lipid bilayer membrane. The primary structure of the cyanobacterial phytoene desaturase is highly conserved with the enzyme from the green alga Dunalliela bardawil (61% identical and 81% similar; [see, Pecker I, Chamovitz D, Mann V, Sandmann G, Boger P and Hirschberg J (1993) Molecular characterization of carotenoid biosynthesis in plants: the phytoene desaturase gene in tomato. In: Murata N (ed) Research in Photosynthesis, Vol III, pp 11-18. Kluwer, Dordrectht]) and from tomato [see, Pecker I, Chamovitz D, Linden H, Sandmann G and Hirschberg J (1992) A single polypeptide catalyzing the conversion of phytoene to ζ -carotene is transcriptionally regulated during tomato fruit ripening. Proc Natl Acad Sci USA 89: 4962-4966], pepper [see, Hugueney P, Romer S, Kuntz M and Camara B (1992) Characterization and molecular cloning of a flavoprotein catalyzing the synthesis of phytofluene and ζ-carotene in Capsicum chromoplasts. Eur J Biochem 209: 399-407] and soybean [see, Bartley GE, Viitanen PV, Pecker I, Chamovitz D, Hirschberg J and Scolnik PA (1991) Molecular cloning and expression in photosynthetic bacteria of a soybean cDNA coding for phytoene desaturase, an enzyme of the carotenoid biosynthesis pathway. Proc Natl Acad Sci USA 88: 6532-6536] (62-65% identical and ~79% similar; [see, Chamovitz D (1993) Molecular analysis of the early steps of carotenoid biosynthesis in cyanobacteria: Phytoene synthase and phytoene 5

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desaturase. Ph.D. Thesis, The Hebrew University of Jerusalem]). The eukaryotic phytoene desaturase polypeptides are larger (64 kDa); however, they are processed during import into the plastids to mature forms whose sizes are comparable to those of the cyanobacterial enzymes.

There is a high degree of structural similarity in carotenoid enzymes of Rhodobacter capsulatus, Erwinia sp. and Neurospora crassa [reviewed in Armstrong GA, Hundle BS and Hearst JE (1993) Evolutionary conservation and similarities of carotenoid biosynthesis gene products structural photosynthetic and nonphotosynthetic organisms. Meth Enzymol 214: 297-311], including in the crtI gene-product, phytoene desaturase. As indicated above, a high degree of conservation of the primary structure of phytoene desaturases also exists among oxygenic photosynthetic organisms. However, there is little sequence similarity, except for the FAD binding sequences at the amino termini, between the "plant-type" crtP gene products and the "bacterial-type" phytoene desaturases (crt1 gene products; 19-23% identities and 42-47% similarities). It has been hypothesized that crtP and crtI are not derived from the same ancestral gene and that they originated independently through convergent evolution [see, Pecker I, Chamovitz D, Linden H, Sandmann G and Hirschberg J (1992) A single polypeptide catalyzing the conversion of phytoene to ζ-carotene is transcriptionally regulated during tomato fruit ripening. Proc Natl Acad Sci USA 89: 4962-4966]. This hypothesis is supported by the different dehydrogenation sequences that are catalyzed by the two types of enzymes and by their different sensitivities to inhibitors.

Although not as definite as in the case of phytoene desaturase, a similar distinction between cyanobacteria and plants on the one hand and other microorganisms is also seen in the structure of phytoene synthase. The crtB gene (formerly psy) encoding phytoene synthase was identified in the genome of Synechococcus sp. strain PCC 7942 adjacent to crtP and within the same operon [see, Bartley GE, Viitanen PV, Pecker I, Chamovitz D, Hirschberg J and Scolnik PA (1991) Molecular cloning and expression in photosynthetic bacteria of a soybean cDNA coding for phytoene desaturase, an enzyme of the carotenoid biosynthesis pathway. Proc Natl Acad Sci USA 88: 6532-6536]. This gene encodes a 36-kDa polypeptide of 307 amino acids with a hydrophobic index of -0.4. The deduced amino acid sequence of the cyanobacterial phytoene synthase is highly conserved with the tomato phytoene synthase (57% identical and 70% similar; Ray JA, Bird CR, Maunders M, Grierson D and Schuch W (1987) Sequence of pTOM5, a ripening related cDNA from tomato. Nucl Acids Res 15: 10587-10588]) but is less highly conserved with the crtB sequences from other

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bacteria (29-32% identical and 48-50% similar with ten gaps in the alignment). Both types of enzymes contain two conserved sequence motifs also found in prenyl transferases from diverse organisms [see, Bartley GE, Viitanen PV, Pecker l, Chamovitz D, Hirschberg J and Scolnik PA (1991) Molecular cloning and expression in photosynthetic bacteria of a soybean cDNA coding for phytoene desaturase, an enzyme of the carotenoid biosynthesis pathway. Proc Natl Acad Sci USA 88: 6532-6536; Carattoli A, Romano N, Ballario P, Morelli G and Macino G (1991) The Neurospora crassa carotenoid biosynthetic gene (albino 3). J Biol Chem 266: 5854-5859; Armstrong GA, Hundle BS and Hearst JE (1993) Evolutionary conservation and structural similarities of carotenoid biosynthesis gene products from photosynthetic and nonphotosynthetic organisms. Meth Enzymol 214: 297-311; Math SK, Hearst JE and Poulter CD (1992) The crtE gene in Erwinia herbicola encodes geranylgeranyl diphosphate synthase. Proc Natl Acad Sci USA 89: 6761-6764; and, Chamovitz D (1993) Molecular analysis of the early steps of carotenoid biosynthesis in cyanobacteria: Phytoene synthase and phytoene desaturase. Ph.D. Thesis, The Hebrew University of Jerusalem]. It is conceivable that these regions in the polypeptide are involved in the binding and/or removal of the pyrophosphate during the condensation of two GGPP molecules.

The crtQ gene encoding ζ -carotene desaturase (formerly zds) was cloned from Anabaena sp. strain PCC 7120 by screening an expression library of cyanobacterial genomic DNA in cells of Escherichia coli carrying the Erwinia sp. crtB and crtE genes and the cyanobacterial crtP gene [see, Linden H, Vioque A and Sandmann G (1993) Isolation of a carotenoid biosynthesis gene coding for ζcarotene desaturase from Anabaena PCC 7120 by heterologous complementation. FEMS Microbiol Lett 106: 99-104]. Since these Escherichia coli cells produce Zcarotene, brownish-red pigmented colonies that produced lycopene could be identified on the yellowish background of cells producing ζ -carotene. predicted ζ-carotene desaturase from Anabaena sp. strain PCC 7120 is a 56-kDa polypeptide which consists of 499 amino acid residues. Surprisingly, its primary structure is not conserved with the "plant-type" (crtP gene product) phytoene desaturases, but it has considerable sequence similarity to the bacterial-type enzyme (crt1 gene product) [see, Sandmann G (1993) Genes and enzymes involved in the desaturation reactions from phytoene to lycopene. (abstract), 10th International Symposium on Carotenoids, Trondheim CL1-2]. It is possible that the cyanobacterial crtQ gene and crtI gene of other microorganisms originated in evolution from a common ancestor.

The crtL gene for lycopene cyclase (formerly lcy) was cloned from Synechococcus sp. strain PCC 7942 utilizing essentially the same cloning strategy

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as for crtP. By using an inhibitor of lycopene cyclase, 2-(4-methylphenoxy)-triethylamine hydrochloride (MPTA), the gene was isolated by transformation of the wild-type to herbicide-resistance [see, Cunningham FX Jr, Chamovitz D, Misawa N, Gantt E and Hirschberg J (1993) Cloning and functional expression in Escherichia coli of a cyanobacterial gene for lycopene cyclase, the enzyme that catalyzes the biosynthesis of β-carotene. FEBS Lett 328: 130-138]. Lycopene cyclase is the product of a single gene product and catalyzes the double cyclization reaction of lycopene to β-carotene. The crtL gene product in Synechococcus sp. strain PCC 7942 is a 46-kDa polypeptide of 411 amino acid residues. It has no sequence similarity to the crtY gene product (lycopene cyclase) from Erwinia uredovora or Erwinia herbicola.

The gene for β-carotene hydroxylase (crtZ) and zeaxanthin glycosilase (crtX) have been cloned from Erwinia herbicola [see, Hundle B, Alberti M, Nievelstein V, Beyer P, Kleinig H, Armstrong GA, Burke DH and Hearst JE (1994) Functional assignment of Erwinia herbicola Eho10 carotenoid genes expressed in Escherichia coli. Mol Gen Genet 254: 406-416; Hundle BS, Obrien DA, Alberti M, Beyer P and Hearst JE (1992) Functional expression of zeaxanthin glucosyltransferase from Erwinia herbicola and a proposed diphosphate binding site. Proc Natl Acad Sci USA 89: 9321-9325] and from Erwinia uredovora [see, Misawa N, Nakagawa M, Kobayashi K, Yamano S, Izawa I, Nakamura K and Harashima K (1990) Elucidation of the Erwinia uredovora carotenoid biosynthetic pathway by functional analysis of gene products in Escherichia coli. J Bacteriol 172: 6704-6712].

The ketocarotenoid astaxanthin (3,3'-dihydroxy-β,β-carotene-4,4'-dione) was first described in aquatic crustaceans as an oxidized form of β-carotene. Astaxanthin was later found to be very common in many marine animals and algae. However, only few animals can synthesize astaxanthin *de novo* from other carotenoids and most of them obtain it in their food. In the plant kingdom, astaxanthin occurs mainly in some species of cyanobacteria, algae and lichens. However, it is found rarely also in petals of higher plant species [see, Goodwin TW (1980) The Biochemistry of the carotenoids, Vol. 1. 2nd Ed, Chapman and Hall, London and New York].

The function of astaxanthin as a powerful antioxidant in animals has been demonstrated [see, Miki W (1991) Biological functions and activities of animal carotenoids. Pure Appl Chem 63: 141]. Astaxanthin is a strong inhibitor of lipid peroxidation and has been shown to play an active role in the protection of biological membranes from oxidative injury [see, Palozza P and Krinsky NI (1992) Antioxidant effects of carotenoids *in vivo* and *in vitro* - an overview. Methods

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Enzymol 213: 403-420; and, Kurashige M, Okimasu E, Inove M and Utsumi K (1990) Inhibition of oxidative injury of biological membranes by astaxanthin. Physiol Chem Phys Med NMR 22: 27]. The chemopreventive effects of astaxanthin have also been investigated in which astaxanthin was shown to significantly reduce the incidence of induced urinary bladder cancer in mice [see, Tanaka T, Morishita Y, Suzui M, Kojima T, Okumura A. and Mori H (1994). Chemoprevention of mouse urinary bladder carcinogenesis by the naturally occurring carotenoid astaxanthin. Carcinogenesis 15: 15]. It has also been demonstrated that astaxanthin exerts immunomodulating effects by enhancing antibody production [see, Jyonouchi H, Zhang L and Tomita Y (1993) Studies of immunomodulating actions of carotenoids. II. Astaxanthin enhances in vitro antibody production to T-dependent antigens without facilitating polyclonal B-cell activation. Nutr Cancer 19: 269; and, Jyonouchi H, Hill JR, Yoshifumi T and Good RA (1991) Studies of immunomodulating actions of carotenoids. I. Effects of β-carotene and astaxanthin on murine lymphocyte functions and cell surface marker expression in-vitro culture system. Nutr Cancer 16: 93]. The complete biomedical properties of astaxanthin remain to be elucidated, but initial results suggest that it could play an important role in cancer and tumor prevention, as well as eliciting a positive response from the immune system.

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Astaxanthin is the principal carotenoid pigment of salmonids and shrimps and imparts attractive pigmentation in the eggs, flesh and skin [see, Torrisen OJ, Hardy RW, Shearer KD (1989) Pigmentation of salmonid-carotenoid deposition and metabolism in salmonids. Crit Rev Aquatic Sci 1: 209]. The world-wide harvest of salmon in 1991 was approximately 720,000 MT., of which 25-30% were produced in a variety of aquaculture facilities [see, Meyers SP (1994) Developments in world aquaculture, feed formulations, and role of carotenoids. Pure Appl Chem 66: 1069]. This is set to increase up to 460,000 MT. by the year 2000 [see, Bjorndahl T (1990) The Economics of Salmon Aquaculture. Blackwell Scientific, Oxford. pp. 1]. The red coloration of the salmonid flesh contributes to consumer appeal and therefore affects the price of the final product. Animals cannot synthesize carotenoids and they acquire the pigments through the food chain from the primary producers - marine algae and phytoplankton. Those grown in intensive culture usually suffer from suboptimal color. Consequently, carotenoid-containing nourishment is artificially added in aquaculture, at considerable cost to the producer.

Astaxanthin is the most expensive commercially used carotenoid compound (todays-1995 market value is of 2,500-3,500 \$/kg). It is utilized mainly as nutritional supplement which provides pigmentation in a wide variety of aquatic

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animals. In the Far-East it is used also for feeding poultry to yield a typical pigmentation of chickens. It is also a desirable and effective nontoxic coloring for the food industry and is valuable in cosmetics. Recently it was reported that astaxanthin is a potent antioxidant in humans and thus is a desirable food additive.

Natural (3S,3'S) astaxanthin is limited in availability. It is commercially extracted from some crustacea species [see, Torrisen OJ, Hardy RW, Shearer KD (1989) Pigmentation of salmonid-carotenoid deposition and metabolism in The (3R,3'R) stereoisomer of Crit Rev Aquatic Sci 1: 209]. salmonids. astaxanthin is produced from Phaffia [a yeast specie, see, Andrewes AG, Phaff HJ and Starr MP (1976) Carotenoids of Phaffia rhodozyma, a red-pigmented fermenting yeast. Phytochemistry Vol. 15, pp. 1003-1007]. Synthetic astaxanthin, comprising a 1:2:1 mixture of the (3S,3'S)-, (3S,3'R)- and (3R,3'R)-isomers is now manufactured by Hoffman-La Roche and sold at a high price (ca. \$2,500/Kg) under the name "CAROPHYLL Pink" [see, Mayer H (1994) Reflections on carotenoid synthesis. Pure & Appl Chem, Vol. 66, pp. 931-938]. Recently a novel gene involved in ketocompound biosynthesis, designated crtW was isolated from the marine bacteria Agrobacterium auranticacum and Alcaligenes PC-1 that produce ketocarotenoids such as astaxanthin. When the crtW gene was introduced into engineered Eschrichia coli that accumulated β-carotene due to Erwinia carotenogenic genes, the Escherichia coli transformants synthesized canthaxanthin a precursor in the synthetic pathway of astaxanthin [see, Misawa N, Kajiwara S, Kondo K, Yokoyama A, Satomi Y, Saito T, Miki W and Ohtani T (1995) Canthaxanthin biosynthesis by the conversion of methylene to keto groups in a hydrocarbon β-carotene by a single gene. Biochemical and biophysical research It is therefore desirable to find a communications Vol. 209, pp. 867-876]. relatively inexpensive source of (3S,3'S) astaxanthin to be used as a feed supplement in aquaculture and as a valuable chemical for various other industrial uses.

Although astaxanthin is synthesized in a variety of bacteria, fungi and algae, the key limitation to the use of biological systems for its production is the low yield of and costly extraction methods in these systems compared to chemical synthesis. One way to solve these problems is to increase the productivity of astaxanthin production in biological systems using recombinant DNA technology. This allows for the production of astaxanthin in genetically engineered host which, in the case of a higher plant, is easy to grow and simple to extract. Furthermore, production of astaxanthin in genetically engineered host enables by appropriate host selection to use thus produced astaxanthin in for example aquaculture applications, devoid of the need for extraction.

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There is thus a widely recognized need for, and it would be highly advantageous to have, a nucleic acid segment which encodes β -C-4-oxygenase, the enzyme that converts β -carotene to canthaxanthin, as well as recombinant vector molecules comprising a nucleic acid sequence according to the invention, and host cells or transgenic organisms transformed or transfected with these vector molecules or DNA segment for the biotechnological production of (3S,3'S) astaxanthin.

Other features and advantages of the invention will be apparent from the following description and from the claims.

SUMMARY OF THE INVENTION

It is a general object of this invention to provide a biotechnological method for production of (3S,3'S) astaxanthin.

It is a specific object of the invention to provide a peptide having a β -C-4-oxygenase activity and a DNA segment coding for this peptide to enable a biotechnological production of astaxanthin and other xanthophylls.

It is a further object of the invention to provide an RNA segments coding for a polypeptide comprising an amino acid sequence corresponding to above described peptide.

It is yet a further object of the invention to provide a recombinant DNA molecule comprising a vector and the DNA segment as described above.

It is still a further object of the invention to provide a host cell containing the above described recombinant DNA molecule.

It is another object of the invention to provide a host transgenic organism containing the above described recombinant DNA molecule or the above described DNA segment in its cells.

It is still another object of the invention to provide a host transgenic organism which expresses β -C-4-oxygenase activity in chloroplasts and/or chromoplasts-containing tissues.

It is yet another object of the invention to provide a food additive for animal or human consumption comprising the above described host cell or transgenic organism.

It is still another object of the invention to provide a method of producing astaxanthin using the above described host cell or transgenic organism.

It is a further object of the invention to provide a method of producing canthaxanthin, echinenone, cryptoxanthin, isocryptoxanthin hydroxyechinenone, zeaxanthin, adonirubin, and/or adonixanthin using the above described host cell or transgenic organism.

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Further objects and advantages of the present invention will be clear from the description that follows.

In one embodiment, the present invention relates to a DNA segment coding for a polypeptide comprising an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene.

In a further embodiment, the present invention relates to an RNA segment coding for a polypeptide comprising an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene.

In yet another embodiment, the present invention relates to a polypeptide comprising an amino acid sequence corresponding to a *Haematococcus pluvialis* crtO gene.

In a further embodiment, the present invention relates to a recombinant DNA molecule comprising a vector and a DNA segment coding for a polypeptide, corresponding to a *Haematococcus pluvialis crtO* gene.

In another embodiment, the present invention relates to a host cell containing the above described recombinant DNA molecule or DNA segment.

In a further embodiment, the present invention relates to a host transgenic organism containing the above described recombinant DNA molecule or the above described DNA segment in its cells.

In another embodiment, the present invention relates to a method of producing astaxanthin using the above described host cell or transgenic organism.

In yet another embodiment, the present invention relates to a method of producing other xanthophylls.

In still another embodiment, the present invention relates to a method of obtaining high expression of a transgene in plants specifically in chromoplasts-containing cells.

In one further embodiment, the present invention relates to a method of importing a carotenoid-biosynthesis enzyme encoded by a transgene into chromoplasts.

BRIEF DESCRIPTION OF THE DRAWINGS

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The invention herein described, by way of example only, with reference to the accompanying drawings, wherein:

FIG. 1 is a general biochemical pathway of β -carotene biosynthesis, in which pathway all molecules are depicted in an all-trans configuration, wherein IPP is isopentenyl pyrophosphate, DMAPP is dimethylallyl pyrophosphate, GPP is geranyl pyrophosphate, FPP is farnesyl pyrophosphate, GGPP is geranyl pyrophosphate and, PPPP is prephytoene pyrophosphate;

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FIG. 2 is an identity map between the nucleotide sequence of the *crtO* cDNA of the present invention (CRTOA.SEQ) and the cDNA cloned by Kajiwara *et al.*, (CRTOJ.SEQ) [see, Kajiwara S, Kakizono T, Saito T, Kondo K, Ohtani T, Nishio N, Nagai S and Misawa N (1995) Isolation and functional identification of a novel cDNA for astaxanthin biosynthesis from *Haematococcus pluvialis*, and astaxanthin synthesis in *Escherichia coli*. Plant Molec Biol 29: 343-352], using a GCG software, wherein (:) indicate identity, (-) indicate a gap and nucleotides numbering is according to SEQ ID NO:4 for CRTOA.AMI and Kajiwara *et al.*, for CRTOJ.AMI;

FIG. 3 is an identity map between the amino acid sequence encoded by the crtO cDNA of the present invention (CRTOA.AMI) and the amino acid sequence encoded by the cDNA cloned by Kajiwara et al., (CRTOJ.AMI) [see, Kajiwara S, Kakizono T, Saito T, Kondo K, Ohtani T, Nishio N, Nagai S and Misawa N (1995) Isolation and functional identification of a novel cDNA for astaxanthin biosynthesis from Haematococcus pluvialis, and astaxanthin synthesis in Escherichia coli. Plant Molec Biol 29: 343-352], using a GCG software, wherein (:) indicate identity, (-) indicate a gap and amino acids numbering is according to SEQ ID NO:4 for CRTOA.AMI and Kajiwara et al., for CRTOJ.AMI;

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FIG. 4 is a schematic depiction of a pACYC184 derived plasmid designated pBCAR and includes the genes crtE, crtB, crtI and crtY of $Erwinia\ herbicola$, which genes are required for production of β -carotene in $Escherichia\ coli$ cells;

FIG. 5 is a schematic depiction of a pACYC184 derived plasmid designated pZEAX and includes the genes *crtE*, *crtB*, *crtI*, *crtY* and *crtZ* from *Erwinia herbicola*, which genes are required for production of zeaxanthin in *Escherichia coli* cells;

FIG. 6 is a schematic depiction of a pBluescriptSK- derived plasmid designated pHPK, containing a full length cDNA insert encoding a β-carotene C-4-oxygenase enzyme from *Haematococcus pluvialis*, designated *crtO* and set forth in SEQ ID NO:1, which cDNA was identified by color complementation of *Escherichia coli* cells;

FIG. 7 is a schematic depiction of a pACYC184 derived plasmid designated pCANTHA which was derived by inserting a 1.2 kb PstI-PstI DNA fragment, containing the cDNA encoding the β-C-4-oxygenase from Haematococcus pluvialis isolated from the plasmid pHPK of Figure 6 and inserted into a PstI site in the coding sequence of the crtZ gene in the plasmid pZEAX of Figure 5; this recombinant plasmid carries the genes crtE, crtB, crtI, crtY of Erwinia herbicola and the crtO gene of Haematococcus pluvialis, all required for production of canthaxanthin in Escherichia coli cells;

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FIG. 8 is a schematic depiction of a pACYC184 derived plasmid designated pASTA which was derived by inserting the 1.2 kb PstI-PstI DNA fragment, containing the cDNA of the β-C-4-oxygenase from Haematococcus pluvialis isolated from the plasmid pHPK of Figure 6 and inserted into a PstI site which exists 600 bp downstream of the crtE gene in the plasmid pZEAX of Figure 5; this recombinant plasmid carries the genes crtE, crtB, crtI, crtY, crtZ of Erwinia herbicola and the crtO gene of Haematococcus pluvialis, all required for production of astaxanthin in Escherichia coli cells;

FIG. 9 is a schematic depiction of a pBR328 derived plasmid designated PAN3.5-KETO which was derived by inserting the 1.2 kb *PstI-PstI* DNA fragment, containing the cDNA of the β-C-4-oxygenase from *Haematococcus pluvialis* isolated from the plasmid pHPK of Figure 6 and inserted into a *PstI* site which exists in a β-lactamase gene in a plasmid designated pPAN35D5 [described in Hirschberg J, Ohad N, Pecker I and Rahat A (1987) Isolation and characterization of herbicide resistant mutants in the cyanobacterium *Synechococcus* R2. Z. Naturforsch 42c: 102-112], which carries the *psbAI* gene from the cyanobacterium *Synechococcus* PCC7942 in the plasmid vector pBR328 [see, Hirschberg J, Ohad N, Pecker I and Rahat A (1987) Isolation and characterization of herbicide resistant mutants in the cyanobacterium *Synechococcus* R2. Z. Naturforsch 42c: 102-112]; this recombinant plasmid carries the *crtO* gene of *Haematococcus pluvialis*, required for production of astaxanthin in *Synechococcus* PCC7942 cells;

FIG. 10 is a schematic depiction of the T-DNA region of a Ti binary plasmid (*E. coli*, *Agrobacterium*) designated pBIB [described by Becker D (1990) Binary vectors which allow the exchange of plant selectable markers and reporter genes. Nucleic Acids Research 18:230] which is a derivative of the Ti plasmid pBI101 [described by Jeffesrson AR, Kavanagh TA and Bevan WM (1987) GUS fusions: β-glucuronidase as a sensitive and versatile gene fusion marker in higher plants. The EMBO J. 6: 3901-3907], wherein BR and BL are the right and left borders, respectively, of the T-DNA region, pAg7 is the polyadenylation site of gene 7 of *Agrobacterium* Ti-plasmid, pAnos is a 250 bp long DNA fragment containing the poly adenylation site of the nopaline synthase gene of *Agrobacterium*, NPT II is a 1,800 bp long DNA fragment containing the promoter sequence of the nopaline synthase gene of *Agrobacterium*, whereas pAnos is a 300 bp long DNA fragment containing the promoter sequence of the nopaline synthase gene of *Agrobacterium*, whereas pAnos is a 300 bp long DNA fragment containing the poly adenylation site of the nopaline synthase gene of *Agrobacterium*;

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FIG. 11 is a schematic depiction of the T-DNA region of a Ti binary plasmid (E. coli, Agrobacterium) designated pPTBIB which was prepared by cloning a genomic DNA sequence of a tomato species Lycopersicon esculentum marked PT (nucleotides 1 to 1448 of the Pds gene as published in Mann V, Pecker I and Hirschberg J (1994) cloning and characterization of the gene for phytoene desaturase (Pds) from tomato (Lycopersicon esculentum). Plant Molecular Biology 24: 429-434), which contains the promoter of the Pds gene and the coding sequence for the amino terminus region of the polypeptide PDS that serve as a transit peptide for import into chloroplasts and chromoplasts, into a HindIII-SmaI site of the binary plasmid vector pBIB of Figure 10, wherein BR and BL, pAg7, pAnos, NPT II, pnos and pAnos are as defined above;

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FIG. 12 is a schematic depiction of the T-DNA region of a Ti binary plasmid (E. coli, Agrobacterium) designated pPTCRTOBIB which was prepared by cloning a 1,110 nucleotide long Eco47III-NcoI fragment of the cDNA of crtO from H. pluvialis (nucleotides 211 to 1321 of SEQ ID NO:1) into the SmaI site of the plasmid pPTBIB of Figure 11, such that the coding nucleotide sequence of the amino terminus of PDS is in the same reading frame of crtO, wherein BR and BL, pAg7, pAnos, NPT II, pnos, and pAnos are as defined above, PT is the promoter and transit peptide coding sequences of Pds from tomato and CRTO is the nucleotide sequence of crtO from H. pluvialis (nucleotides 211 to 1321 of SEQ ID NO:1);

FIG. 13 shows a Southern DNA blot analysis of *Hind*III-digested genomic DNA extracted from wild type (WT) and *crtO* tobacco transgenic plants, designated 2, 3, 4, 6, 9 and 10, according to the present invention, using the *crtO* cDNA as a radioactive probe essentially as described in Sambrook *et al.*, Molecular Cloning; A Laboratory Manual. Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y. 1989, wherein the size of marker (M) DNA fragments in kilobase pairs (kb) is indicated on the left as well as the expected position (arrow) of an internal T-DNA *Hind*III fragment as was deduced from the sequence of pPTPDSBIB shown in Figure 12 which contain the *crtO* cDNA sequence;

FIG. 14 shows a biosynthesis pathway of astaxanthin;

FIG. 15 shows a flower from a wild type tobacco plant and a flower from a transgenic tobacco plant according to the present invention.

DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention is, in general, of a biotechnological method for production of (3S,3'S) astaxanthin. In particular, the present invention is of a

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peptide having a β -C-4-oxygenase activity; a DNA segment coding for this peptide; an RNA segments coding for this peptide; a recombinant DNA molecule comprising a vector and the DNA segment; a host cell or organism containing the above described recombinant DNA molecule or DNA segment; and of a method for biotechnologically producing (3S,3'S) astaxanthin or a food additive containing (3S,3'S) astaxanthin, using the host.

alga Haematococcus pluvialis green The unicellular fresh-water accumulates large amounts of (3S,3'S) astaxanthin when exposed to unfavorable growth conditions, or following different environmental stresses such as phosphate or nitrogen starvation, high concentration of salt in the growth medium or high light intensity [see, Yong YYR and Lee YK (1991) Phycologia 30 257-261; Droop MR (1954) Arch Microbiol 20: 391-397; and, Andrewes A.G, Borch G, Liaaen-Jensen S and Snatzke G.(1974) Acta Chem Scand B28: 730-736]. During this process, the vegetative cells of the alga form cysts and change their color from green to red. The present invention discloses the cloning of a cDNA from Haematococcus pluvialis, designated crtO, which encodes a β-C-4-oxygenase, the enzyme that converts β-carotene to canthaxanthin, and its expression in a heterologous systems expressing β-carotene hydroxylase (e.g., Erwinia herbicola crtZ gene product), leading to the production of (3S,3'S) astaxanthin.

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The crtO cDNA and its encoded peptide having a β -C-4-oxygenase activity are novel nucleic and amino acid sequences, respectively. The cloning method of the crtO cDNA took advantage of a strain of $Escherichia\ coli$, which was genetically engineered to produce β -carotene, to which a cDNA library of $Haematococcus\ pluvialis$ was transfected and expressed. Visual screening for brown-red pigmented $Escherichia\ coli$ cells has identified a canthaxanthin producing transformant. Thus cloned cDNA has been expressed in two heterologous systems ($Escherichia\ coli$ and $Synechococcus\ PCC7942\ cells$) both able to produce β -carotene and further include an engineered ($Erwinia\ herbicola\ crtZ$ gene product) or endogenous β -carotene hydroxylase activity, and was shown to enable the production of (3S,3'S) astaxanthin in both these systems.

The crtO cDNA or its protein product exhibit no meaningful nucleic- or amino acid sequence similarities to the nucleic- or amino acid sequence of crtW and its protein product isolated from the marine bacteria Agrobacterium auranticacum and Alcaligenes PC-1 that produce ketocarotenoids such as astaxanthin [see, Misawa N, Kajiwara S, Kondo K, Yokoyama A, Satomi Y, Saito T, Miki W and Ohtani T (1995) Canthaxanthin biosynthesis by the conversion of methylene to keto groups in a hydrocarbon β-carotene by a single gene. Biochemical and biophysical research communications Vol. 209, pp. 867-876].

However, the *crtO* cDNA and its protein product exhibit substantial nucleic- and amino acid sequence identities with the nucleic- and amino acid sequence of a recently cloned cDNA encoding a 320 amino acids protein product having β-carotene oxygenase activity, isolated from *Haematococcus pluvialis* [see, Kajiwara S, Kakizono T, Saito T, Kondo K, Ohtani T, Nishio N, Nagai S and Misawa N (1995) Isolation and functional identification of a novel cDNA for astaxanthin biosynthesis from *Haematococcus pluvialis*, and astaxanthin synthesis in *Escherichia coli*. Plant Molec Biol 29: 343-352]. Nevertheless, as presented in Figure 2 the degree of sequence identity between the *crtO* cDNA (CRTOA.SEQ in Figure 2) and the cDNA described by Kajiwara *et al.* (CRTOJ.SEQ in Figure 2) [see reference above] is 75.7% and, as presented in Figure 3 the degree of sequence identity between the *crtO* cDNA protein product (CRTOA.AMI in Figure 3) and the protein described by Kajiwara *et al.* (CRTOJ.AMI in Figure 3) is 78%, as was determined using a GCG software.

As will be described in details hereinbelow, the *crtO* cDNA can thus be employed to biotechnologically produce (3S,3'S) astaxanthin in systems which are either easy to grow and can be used directly as an additive to fish food, or systems permitting a simple and low cost extraction procedure of astaxanthin.

In one embodiment, the present invention relates to a DNA segment coding for a polypeptide comprising an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene and allelic and species variations and functional naturally occurring and/or man-induced variants thereof. The phrase 'allelic and species variations and functional naturally occurring and/or man-induced variants' as used herein and in the claims below refer to the source of the DNA (or RNA as described below) or means known in the art for obtaining it. However the terms 'variation' and 'variants' indicate the presence of sequence dissimilarities (i.e., variations). It is the intention herein and in the claims below that the sequence variations will be 77-80%, preferably 80-85%, more preferably 85-90%, most preferably 90-100% of identical nucleotides. In a preferred embodiment the DNA segment comprises the sequence set forth in SEQ ID NO:1. In another preferred embodiment, the DNA segment encodes the amino acid sequence set forth in SEQ ID NO:4.

The invention also includes a pure DNA segment characterized as including a sequence which hybridizes under high stringency conditions [e.g., as described in Sambrook et al., Molecular Cloning; A Laboratory Manual. Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y. 1989] to a nucleic acid probe which includes at least fifteen, preferably at least fifty, more preferably at least hundred, even more preferably at least five

hundred successive nucleotides of SEQ ID NO:1 or SEQ ID NO:2. Alternatively, the DNA segment of the invention may be characterized as being capable of hybridizing under low-stringent conditions to a nucleic acid probe which includes the coding sequence (nucleotides 166 through 1152) of SEQ ID NO:1 or SEQ ID NO:2. An example of such low-stringency conditions is as described in Sambrook et al., using a lower hybridization temperature, such as, for example, 20°C below the temperature employed for high-stringency hybridization conditions, as described above.

The DNA segment of the invention may also be characterized as being capable of hybridizing under high-stringent conditions to a nucleic acid probe which includes the coding sequence (nucleotides 166 through 1152) of SEQ ID NO:1 or SEQ ID NO:2.

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The invention also includes a synthetically produced oligonucleotide (e.g., oligodeoxyribonucleotide or oligoribonucleotide and analogs thereof) capable of hybridizing with at least ten-nucleotide segments of SEQ ID NO:1 or SEQ ID NO:2.

In another embodiment, the present invention relates to an RNA segment coding for a polypeptide comprising an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene and allelic and species variations and functional naturally occurring and/or man-induced variants thereof. In a preferred embodiment the RNA segment comprises the sequence set forth in SEQ ID NO:2. In another preferred embodiment, the RNA segment encodes the amino acid sequence set forth in SEQ ID NO:4.

The invention also includes a pure RNA characterized as including a sequence which hybridizes under high stringent conditions to a nucleic acid probe which includes at least at least fifteen, preferably at least fifty, more preferably at least hundred, even more preferably at least five hundred successive nucleotides of SEQ ID NO:1 or SEQ ID NO:2. Alternatively, the RNA of the invention may be characterized as being capable of hybridizing under low-stringent conditions to a nucleic acid probe which includes the coding sequence (nucleotides 166 through 1152) of SEQ ID NO:1 or SEQ ID NO:2. Additionally, the RNA of the invention may be characterized as being capable of hybridizing under high-stringent conditions to a nucleic acid probe which includes the coding sequence (nucleotides 166 through 1152) of SEQ ID NO:1 or SEQ ID NO:2.

In another embodiment, the present invention relates to a polypeptide comprising an amino acid sequence corresponding to a *Haematococcus pluvialis* crtO gene and allelic, species variations and functional naturally occurring and/or

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man-induced variants thereof.

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In a preferred embodiment, the polypeptide comprises the amino acid sequence set forth in SEQ ID NO:4.

It should be noted that the invention includes any peptide which is homologous (i.e., 80-85%, preferably 85-90%, more preferably 90-100% of identical amino acids) to the above described polypeptide. The term 'homologous' as used herein and in the claims below, refers to the sequence identity between two peptides. When a position in both of the two compared sequences is occupied by identical amino acid monomeric subunits, it is homologous at that position. The homology between two sequences is a function of the number of homologous positions shared by the two sequences. For example, if eight of ten of the positions in two sequences are occupied by identical amino acids then the two sequences are 80% homologous.

Other polypeptides which are also included in the present invention are allelic variations, other species homologs, natural mutants, induced mutants and peptides encoded by DNA that hybridizes under high or low stringency conditions (see above) to the coding region (nucleotides 166 through 1152) of SEQ ID NO:1 or SEQ ID NO:2.

In another embodiment, the present invention relates to a recombinant DNA molecule comprising a vector (for example plasmid or viral vector) and a DNA segment coding for a polypeptide, as described above. In a preferred embodiment, the DNA segment is present in the vector operably linked to a promoter.

In a further embodiment, the present invention relates to a host cell containing the above described recombinant DNA molecule or DNA segment. Suitable host cells include prokaryotes (such as bacteria, including Escherichia coli) and both lower eukaryotes (for example yeast) and higher eukaryotes (for example, algae, plant or animal cells). Introduction of the recombinant molecule into the cell can be effected using methods known in the art such as, but not limited to, transfection, transformation, micro-injection, gene bombardment etc. The cell thus made to contain the above described recombinant DNA molecules may be grown to form colonies or may be made to differentiate to form a differentiated organism. The recombinant DNA molecule may be transiently contained (e.g., by a process known in the art as transient transfection) in the cell, nevertheless, it is preferred that the recombinant DNA molecule is stably contained (e.g., by a process known in the art as stable transfection) in the cell. Yet in a preferred embodiment the cell is endogenously producing, or is made by genetic engineering means to produce, β-carotene, and the cell contains endogenous or genetically engineered β-carotene hydroxylase activity. Such a cell may be used as a food additive for animal (e.g., salmon) and human consumption. Furthermore.

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such a cell may be used for extracting astaxanthin and/or other xanthophylls, as described hereinbelow.

In a further embodiment, the present invention relates to a host transgenic organism (e.g., a higher plant or animal) containing the above described recombinant DNA molecule or the above described DNA segment in its cells. Introduction of the recombinant molecule or the DNA segment into the host transgenic organism can be effected using methods known in the art. Yet, in a preferred embodiment the host organism is endogenously producing, or is made by genetic engineering means to produce, β -carotene and, also preferably the host organism contains endogenous or genetically engineered β -carotene hydroxylase activity. Such an organism may be used as a food additive for animal (e.g., salmon) and human consumption. Furthermore, such an organism may be used for extracting astaxanthin and/or other xanthophylls, as described hereinbelow.

In another embodiment, the present invention relates to a method of producing astaxanthin using the above described host cell or transgenic organism. In yet another embodiment, the present invention relates to a method of producing xanthophylls such as canthaxanthin, echinenone, cryptoxanthin, isocryptoxanthin, hydroxyechinenone, zeaxanthin, adonirubin, 3-hydroxyechinenone, 3'-hydroxyechinenone and/or adonixanthin using the above described host cell or transgenic organism. For these purposes provided is a cell or a transgenic organism as described above. The host cell or organism are made to grow under conditions favorable of producing astaxanthin and the above listed additional xanthophylls which are than extracted by methods known in the art.

In yet another embodiment, the present invention relates to a transgenic plant expressing a transgene coding for a polypeptide including an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene, allelic and species variants or functional naturally occurring or man-induced variants thereof. Preferably the expression is highest in chromoplasts-containing tissues.

In yet another embodiment, the present invention relates to a recombinant DNA vector which includes a first DNA segment encoding a polypeptide for directing a protein into plant chloroplasts or chromoplasts (e.g., derived from the Pds gene of tomato) and an in frame second DNA segment encoding a polypeptide including an amino acid sequence corresponding to Haematococcus pluvialis crtO gene, allelic and species variants or functional naturally occurring and maninduced variants thereof.

In yet another embodiment, the present invention relates to a recombinant DNA vector which includes a first DNA segment including a promoter highly expressible in plant chloroplasts or chromoplasts-containing tissues (e.g., derived

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from the *Pds* gene of tomato) and a second DNA segment encoding a polypeptide including an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene, allelic and species variants or functional naturally occurring and maninduced variants thereof.

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Reference in now made to the following examples, which together with the above descriptions, illustrate the invention.

EXAMPLES

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The following protocols and experimental details are referenced in the Examples that follow:

Algae and growth conditions. Haematococcus pluvialis (strain 34/7 from the Culture Collection of Algae and Protozoa, Windermere, UK) was kindly provided by Dr. Andrew Young from the Liverpool John Moores University. Suspension cultures of the alga were grown in a liquid medium as described by Nichols and Bold [see, Nichols HW, Bold HC (1964) Trichsarcina polymorpha gen et sp nov J Phycol 1: 34-39]. For induction of astaxanthin biosynthesis cells were harvested, washed in water and resuspended in a nitrogen-depleted medium. The cultures were maintained in 250 ml Erlenmeyer flasks under continuous light (photon flux of 75 μ E/m²/s), at 25°C, on a rotary shaker at 80 rpm.

Construction of cDNA library. The construction of a cDNA library from Haematococcus pluvialis was described in detail by Lotan and Hirschberg (1995) FEBS letters 364: 125-128. Briefly, total RNA was extracted from algal cells grown for 5 days under nitrogen-depleted conditions (cell color brown-red). Cells from a 50 ml culture were harvested and their RNA content was extracted using Tri reagent (Molecular Research Center, INC.). Poly-An RNA was isolated by two cycles of fractionation on oligo dT-cellulose (Boehringer). The final yield was 1.5% of the total RNA. The cDNA library was constructed in a Uni-ZAPTM XR vector, using a ZAP-cDNA synthesis kit (both from Stratagene). Escherichia coli cells of strain XL1-Blue MRF' (Stratagene) were used for amplification of the cDNA library.

Plasmids and Escherichia coli strains. Plasmid pPL376, which contains the genes necessary for carotenoid biosynthesis in the bacterium Erwinia herbicola was obtained from Tuveson [for further details regarding plasmid pPL376 see, Tuveson RW, Larson RA & Kagan J (1988) Role of cloned carotenoid genes expressed in Escherichia coli in protecting against inactivation by near-UV light

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and specific phototoxic molecules. J Bacteriol 170: 4675-4680]. Cells of Escherichia coli strain JM109 that carry the plasmid pPL376 accumulate the bright yellow carotenoid, zeaxanthin glycoside. In a first step, a 1.1 kb SalI-SalI fragment was deleted from this plasmid to inactivate the gene crtX, coding for zeaxanthin glucosyl transferase. In a second step, partial BamHI cleavage of the plasmid DNA, followed by self ligation, deleted a 0.8 kb fragment which inactivated crtZ, encoding β-carotene hydroxylase. A partial BglII cleavage generated a fragment of 7.4 kb which was cloned in the BamHI site of the plasmid vector pACYC184. As shown in Figure 4, the resulting recombinant plasmid, which carried the genes crtE, crtB, crtI and crtY, was designated pBCAR [Lotan and Hirschberg (1995) FEBS letters 364: 125-128].

Plasmid pBCAR was transfected into SOLR strain cells of *Escherichia coli* (Stratagene). Colonies that appeared on chloramphenicol-containing Luria Broth (LB) medium [described in Sambrook *et al.*, Molecular Cloning; A Laboratory Manual. Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y. 1989], carried this plasmid and developed a deep yellow-orange color due to the accumulation of β-carotene.

As shown in Figure 5, an additional plasmid, designated pZEAX, which allows for zeaxanthin synthesis and accumulation in *Escherichia coli* was constructed [this plasmid is described in details in Lotan and Hirschberg (1995) FEBS letters 364: 125-128]. SOLR strain *Escherichia coli* cells were used as a host for the pZEAX plasmid. *Escherichia coli* cells were grown on LB medium (see above), at 37°C in the dark on a rotary shaker at 225 rpm. Ampicillin (50 μ g/ml) and/or chloramphenicol (30 μ g/ml) (both from Sigma) were added to the medium for selection of appropriate transformed cells.

As shown in Figure 6, a plasmid, pHPK, containing the full length cDNA of the β-carotene C-4-oxygenase enzyme was identified by color complementation as described by Lotan and Hirschberg (1995) FEBS letters 364: 125-128 (see description herein below). A 1.2 kb *PstI-PstI* DNA fragment, containing the cDNA of the β-C-4-oxygenase from *Haematococcus pluvialis*, was isolated from plasmid pHPK and inserted into a *PstI* site in the coding sequence of the *crtZ* gene in the plasmid pZEAX. This recombinant plasmid was designated pCANTHA and is shown in Figure 7.

The same 1.2 kb *PstI-PstI* fragment was also inserted into a *PstI* site which exists 600 bp downstream of the *crtE* gene in the plasmid pZEAX. The resulting recombinant plasmid was designated pASTA and is shown in Figure 8.

The same 1.2 kb PstI-PstI fragment was also inserted into a PstI site which exists in the β -lactamase gene in the plasmid pPAN35D5 [Hirschberg J, Ohad N,

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Pecker I and Rahat A (1987) Isolation and characterization of herbicide resistant mutants in the cyanobacterium *Synechococcus* R2. Z. Naturforsch 42c: 102-112], which carries the *psbAI* gene from the cyanobacterium *Synechococcus* PCC7942 in the plasmid vector pBR328 [Hirschberg J, Ohad N, Pecker I and Rahat A (1987) Isolation and characterization of herbicide resistant mutants in the cyanobacterium *Synechococcus* R2. Z. Naturforsch 42c: 102-112]. This plasmid was designated PAN3.5-KETO and is shown in Figure 9. This plasmid was used in the transformation of *Synechococcus* PCC7942 cells following procedures described by Golden [Golden SS (1988) Mutagenesis of cyanobacteria by classical and gene-transfer-based methods. Methods Enzymol 167: 714-727].

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Excision of phage library and screening for a β -carotene oxygenase gene. Mass excision of the cDNA library, which was prepared as described hereinabove, was carried out using the ExAssist helper phage (Stratagene) in cells of SOLR strain of *Escherichia coli* that carried the plasmid pBCAR. The excised library in phagemids form was transfected into *Escherichia coli* cells strain XL1-Blue and the cells were plated on LB plates containing 1 mM isopropylthio- β -D-galactosidase (IPTG), 50 μ g/ml ampicillin and 30 μ g/ml chloramphenicol, in a density that yielded approximately 100-150 colonies per plate. The plates were incubated at 37°C overnight and further incubated for two more days at room temperature. The plates were then kept at 4°C until screened for changes in colony colors.

A plasmid for high expression of crtO in chromoplasts. As shown in Figures 10-11, a genomic DNA sequence of a tomato species Lycopersicon esculentum (nucleotides 1 to 1448 of the Pds gene [as published in Mann V, Pecker I and Hirschberg J (1994) cloning and characterization of the gene for phytoene desaturase (Pds) from tomato (Lycopersicon esculentum). Plant Molecular Biology 24: 429-434], which contains the promoter of the Pds gene and the coding sequence for the amino terminus region of the polypeptide PDS that serve as a transit peptide for import into chloroplasts and chromoplasts, was cloned into a HindIII-SmaI site of the binary plasmid vector pBIB, [described by Becker D (1990) Binary vectors which allow the exchange of plant selectable markers and reporter genes. Nucleic Acids Research 18:230], shown in Figure 10. The recombinant plasmid was designated pPTBIB and is shown in Figure 11.

As shown in Figure 12, a 1,110 nucleotide long *Eco*47III-*Nco*I fragment, containing the cDNA of *crtO* from *H. pluvialis* (nucleotides 211 to 1321 of SEQ ID NO:1) was sub-cloned into the *Sma*I site of the plasmid pPTBIB (Figure 11) so that the coding nucleotide sequence of the amino terminus of

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Pds is in the same reading frame as crtO. The recombinant plasmid was designate pPTCRTOBIB.

Formation of transgenic higher plant. The DNA of pPTCRTOBIB was extracted from E. coli cells and was transferred into cells of Agrobacterium tumefaciens strain EHA105 [described by Hood EE, Gelvin SB, Melchers LS and Hoekema A (1993) Transgenic Research 2:208-218] using electroporation as described for E. coli [Dower JW, Miller FJ and Ragdsale WC (1988) High efficiency transformation of E. coli by high voltage electroporation. Nuc. Acids Res. 18: 6127-6145]. Agrobacterium cells were grown at 28 °C in LB medium supplemented with 50 µg/ml streptomycin and 50 µg/ml kanamycin as selective agents. Cells of Agrobacterium carrying pPTCRTOBIB were harvested from a suspension culture at the stationary phase of growth and used for transformation as described by Horsch RB, Fry JE, Hoffmann NL, Eicholtz D, Rogers SG and Fraley RT, A simple and general method for transferring genes into plants. Science (1985) 227:1229-1231; and Jeffesrson AR, Kavanagh TA and Bevan WM (1987) GUS fusions: βglucuronidase as a sensitive and versatile gene fusion marker in higher plants. The EMBO J. 6: 3901-3907.

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Leaf explants of *Nicotiana tobaccum* strain NN were infected with the transformed *Agrobacterium* cells and kanamycin-resistant transgenic plants were regenerated according to protocols described by Horsch et al. (1985) and Jefferson et al. (1987) cited above.

With reference now to Figure 13, the presence of the DNA sequence of the crtO gene-construct in the fully developed regenerated plants was determined by DNA Southern blot analysis. To this end DNA was extracted from the leaves [according to a protocol described by Kanazawa and Tsutsumi (1992) Extraction of restrictable DNA from plants of the genus Nelumbo. Plant Molecular Biology Reports 10: 316-318], digested with the endonuclease HindIII, the fragments were size separated by gel electrophoresis and hybridized with radioactively labeled crtO sequence (SEQ ID NO: 1).

It was determined that each transgenic plant that was examined contained at least one copy of the *crtO* DNA sequence, yielding a 1.75 kb band (arrow), originating from an internal *Hind*III-*Hind*III fragment of the T-DNA of pPTCRTOBIB, additional bands originating from partial digestion, additional band/s whose sizes vary, depending on the position of insertion in the plant genome and a 1.0 kb band originating from the tobacco plant itself which therefore also appears in the negative control WT lane.

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Sequence analysis. DNA sequence analysis was carried out by the dideoxy method [see, Sanger F, Nicklen S & Coulsen AR (1977) DNA sequencing with chain termination inhibitors. Proc Natl Acad Sci USA 74: 5463-5467].

Carotenoids analysis. Aliquots of *Escherichia coli* cells which were grown in liquid in LB medium were centrifuged at 13,000 g for 10 minutes, washed once in water and re-centrifuged. After removing the water the cells were resuspended in 70 μ l of acetone and incubated at 65°C for 15 minutes. The samples were centrifuged again at 13,000 g for 10 minutes and the carotenoid-containing supernatant was placed in a clean tube. The carotenoid extract was blown to dryness under a stream of nitrogen (N₂) gas and stored at -20°C until required for analysis. Carotenoids from plant tissues were extracted by mixing 0.5-1.0 gr of tissue with 100 μ l of acetone followed by incubation at 65°C for 15 minutes and then treating the samples as described above.

High-performance liquid chromatography (HPLC) of the carotenoid extracts was carried out using an acidified reverse-phase C18 column, Spherisorb ODS-2 (silica 5 μ m 4.6 mm x 250 mm) (Phenomenex®). The mobile phase was pumped by triphasic Merck-Hitachi L-6200A high pressure pumps at a flow rate of 1.5 ml/min. The mobile phase consisted of an isocratic solvent system comprised of hexane/dichloromethane/isopropyl alcohol/triethylamine (88.5:10:1.5:0.1, v/v). Peaks were detected at 470 nm using a Waters 996 photodiode-array detector. Individual carotenoids were identified by their retention times and their typical absorption spectra, as compared to standard samples of chemically pure β -carotene, zeaxanthin, echinenone, canthaxanthin, adonirubin and astaxanthin (The latter four were kindly provided by Dr. Andrew Young from Liverpool John Moores University).

Thin layer chromatography (TLC) was carried out using silica gel 60 F254 plates (Merck), using ethyl acetate/benzene (7:3, v/v) as an eluent. Visible absorption spectra were recorded with a Shimadzu UV-160A spectrophotometer. All spectra were recorded in acetone. Spectral fine structure was expressed in terms of %III/II [Britton, G. (1995). UV/Visible Spectroscopy. In: Carotenoids; Vol IB, Spectroscopy. Eds. Britton G, Liaaen-Jensen S and Pfander H. Birkhauser Verlag, Basel. pp. 13-62].

Isolation and identification of the carotenoids extracted from cells of E. coli are treated in order of increasing adsorption (decreasing R_f values) on silica TLC plates. Carotenoids structure and the biosynthesis pathway of astaxanthin are given in Figure 14. The following details refer to the carotenoids numbered 1 through 9 in Figure 14.

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β-Carotene (1). R_f 0.92 inseparable from authentic (1). R_t .VIS λ_{max} nm: (428), 452, 457, %III/II = 0.

Echinenone (2). R_f 0.90 inseparable from authentic (2). R_t .VIS λ_{max} nm: 455. %III/II = 0.

Canthaxanthin (3). R_f 0.87. inseparable from authentic (3). R_t .VIS λ_{max} nm: 470. %III/II = 0.

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β-Cryptoxanthin (4). R_f 0.83. R_t .VIS λ_{\max} nm: (428), 451, 479, %III/II = 0.

Adonirubin (5). R_f 0.82 inseparable from authentic (5). R_t .VIS λ_{max} nm: 476, %III/II = 0.

Astaxanthin (6). R_f 0.79 inseparable from authentic (6). R_t .VIS λ max nm: 477, %III/II = 0.

Adonixanthin (7). $R_f 0.72$. R_t .VIS λ_{max} nm: 464, %III/II = 0.

Zeaxanthin (8). R_f 0.65 inseparable from authentic (8). R_t .VIS λ_{max} nm: (428). 451, 483, %III/II = 27.

Hydroxyechinenone (9). Rf 0.80, Rt, 3.0. VIS λ_{max} nm: 464, %III/II = 0.

Chirality configuration. Chirality configuration of astaxanthin was determined by HPLC of the derived diastereoisomeric camphanates of the astaxanthin [Renstrom B, Borch G, Skulberg M and Liaaen-Jensen S (1981) Optical purity of (3S,3S)-astaxanthin from *Haematococcus pluvialis*. Phytochem 20: 2561-2565]. The analysis proved that the *Escherichia coli* cells synthesize pure (3S,3'S) astaxanthin.

EXAMPLE 1 Cloning the β-C-4-oxygenase gene

A cDNA library was constructed in Lambda ZAP II vector from poly-An RNA of Haematococcus pluvialis cells that had been induced to synthesize astaxanthin by nitrogen deprivation as described hereinabove. The entire library was excised into β -carotene-accumulating cells of Escherichia coli, strain SOLR, which carried plasmid pBCAR (shown in Figure 4). Screening for a β -carotene oxygenase gene was based on color visualization of colonies of size of 3 mm in diameter. Astaxanthin and other oxygenated forms of β -carotene (i.e., xanthophylls) have distinct darker colors and thus can be detected from the yellow β -carotene background. The screening included approximately 100,000 colonies which were grown on LB medium plates containing ampicillin and chloramphenicol that selected for both the Lambda ZAP II vector in its plasmid propagating form and the pBCAR plasmid. Several colonies showed different

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color tones but only one exhibited a conspicuous brown-red pigment. This colony presumed to contain a xanthophyll biosynthesis gene was selected for further analysis described hereinbelow in the following Examples.

EXAMPLE 2

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Analysis of the β -C-4-oxygenase activity in Escherichia coli

The red-brown colony presumed to contain a xanthophyll biosynthesis gene (see Example 1 above) was streaked and further analyzed. First, the recombinant ZAP II plasmid carrying the cDNA clone that was responsible for xanthophyll synthesis in *Escherichia coli* was isolated by preparing plasmid DNA from the red-brown colony, transfecting it to *Escherichia coli* cells of the strain XL1-Blue and selection on ampicillin-containing medium. This plasmid, designated pHPK (pHPK is a Lambda ZAP II vector containing an insert isolated from the red-brown colony), was used to transform β-carotene-producing *Escherichia coli* cells (*Escherichia coli* SOLR strain that carry the plasmid pBCAR shown in Figure 4) resulting in the formation of red-brown colonies. Carotenoids from this transformant, as well as from the host cells (as control) were extracted by acetone and analyzed by HPLC.

HPLC analysis of carotenoids of the host bacteria which synthesized β -carotene (*Escherichia coli* SOLR strain that carry the plasmid pBCAR shown in Figure 4), as compared with a brown-red colony, revealed that only traces of β -carotene were observed in the transformant cells while a new major peak of canthaxanthin and another minor peak of echinenone appeared [described in detail by Lotan and Hirschberg (1995) FEBS letters 364: 125-128]. These results indicate that the cDNA in plasmid pHPK, designated *crtO* encodes an enzyme with β -C-4-oxygenase activity, which converts β -carotene to canthaxanthin *via* echinenone (see Figure 14). It is, therefore concluded that a single enzyme catalyzes this two-step ketonization conversion by acting symmetrically on the 4 and 4' carbons of the β - and β '-rings of β -carotene, respectively.

EXAMPLE 3

Production of astaxanthin in Escherichia coli cells

To determine whether β -carotene hydroxylase (e.g., a product of the crtZ gene of $Erwinia\ herbicola$) can convert thus produced canthaxanthin to astaxanthin and/or whether zeaxanthin converted from β -carotene by β -carotene hydroxylase can be converted by β -C-4-oxygenase to astaxanthin, the crtO cDNA of

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Haematococcus pluvialis thus isolated, was expressed in Escherichia coli cells together with the crtZ gene of Erwinia herbicola. For this purpose, Escherichia coli cells of strain SOLR were transfected with either plasmid pASTA alone containing, as shown in Figure 8, both crtZ and crtO or, alternatively with both plasmids, pHPK containing, as shown in Figure 6, crtO, and pZEAX containing, as shown in Figure 5, crtZ. Carotenoids in the resulting transformed cells were extracted and analyzed by HPLC as described above. The results, given in Table 1, show the composition of carotenoids extracted from the cells containing the plasmid pASTA. Similar carotenoid composition is found in Escherichia coli cells which carry both pHPK and pZEAX.

TABLE 1

	Carotenoid	% of total carotenoid composition
15	•	
	β-Carotene	8.0
	Echineone	1.7
	β-Cryptoxanthin	4.2
	Canthaxanthin	4.2
20	Zeaxanthin	57.8
	Adonirubin	1.0
	Adonixanthin	17.9
	Astaxanthin	5.2

The results presented in Table 1, prove that carotenoids possessing either a β-end group or a 4-keto-β-end group act as substrates for the hydroxylation reactions catalyzed by crtZ gene product at carbons C-3 and C-3'. hydroxylation of β -carotene and canthaxanthin results in the production of zeaxanthin and astaxanthin, respectively. These hydroxylations result in the intermediate ketocarotenoids, production of astaxanthin and the hydroxyechinenone, adonixanthin and adonirubin. These results further demonstrate that astaxanthin can be produced in heterologous cells by expressing the gene crtO together with a gene that codes for a β -carotene hydroxylase.

EXAMPLE 4 Sequence analysis of the gene for β -carotene C-4-oxygenase

The full length, as was determined by the presence of a poly A tail, of the cDNA insert in plasmid pHPK (1771 base pairs) was subjected to nucleotide

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sequence analysis. This sequence, set forth in SEQ ID NO:1, and its translation to an amino acid sequence set forth in SEQ ID NO:3 (329 amino acids), were deposited in EMBL database on May 1, 1995, and obtained the EMBL accession numbers X86782 and X86783, respectively.

An open reading frame (ORF) of 825 nucleotides (nucleotides 166 through 1152 in SEQ ID NO:3) was identified in this sequence. This ORF codes for the enzyme β -carotene C-4-oxygenase having 329 amino acids set forth in SEQ ID NO:4, as proven by its functional expression in *Escherichia coli* cells (see Example 3 above). The gene for this enzyme was designated crtO.

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EXAMPLE 5 Transformation of cyanobacteria with crtO

The plasmid DNA of pPAN3.5-KETO, shown in Figure 9, was transfected into cells of the cyanobacterium *Synechococcus* PCC7942 according to the method described by Golden [Golden SS (1988) Mutagenesis of cyanobacteria by classical and gene-transfer-based methods. Methods Enzymol 167: 714-727]. The cyanobacterial cells were plated on BG11 medium-containing petri dishes that contained also chloramphenicol. Colonies of chloramphenicol-resistant *Synechococcus* PCC7942 which appeared after ten days were analyzed for their carotenoid content. As detailed in Table 2 below, HPLC analysis of these cells revealed that the major carotenoid components of the cells was β-carotene, echinenone, canthaxanthin, adonirubin and astaxanthin. A similar analysis of the wild type strain and of *Synechococcus* PCC7942 transfected with a plasmid in which the orientation of the *crtO* gene is reversed (not shown), which is therefore not capable of producing an active protein, did not revealed production of echinenone, canthaxanthin, adonirubin and astaxanthin.

These result prove that crtO of Haematococcus pluvialis can be expressed in cyanobacteria and that its expression provided a β-C-4-oxygenase enzymatic activity needed for the conversion of \beta-carotene to canthaxanthin. This result endogenous further demonstrates that the β-carotene hydroxylase Synechococcus PCC7942 is able to convert thus produced canthaxanthin to astaxanthin. Since the carotenoid biosynthesis pathway is similar in all green photosynthetic organism [see Figures 1 and 10 and, Pecker I, Chamovitz D. Linden H, Sandmann G and Hirschberg J (1992) A single polypeptide catalyzing the conversion of phytoene to ζ -carotene is transcriptionally regulated during tomato fruit ripening. Proc Natl Acad Sci USA 89: 4962-4966] it is deduced that astaxanthin can be produced in algae, and higher plants by expressing crtO in any

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tissue that express also the endogenous β -carotene hydroxylase. It is further deduced that astaxanthin can be produced by any organism provided it contains either endogenous or engineered β -carotene biosynthesis pathway, by expressing crtO in any tissue that express either endogenous or genetically engineered β -carotene hydroxylase.

TABLE 2

	Carotenoid	% of total carotenoid composition					
10							
	β-Carotene	31.5					
	Echinenone	18.5					
	Canthaxanthin	16.1					
	Zeaxanthin	22.3					
15	Adonirubin	6.0					
	Astaxanthin	5.6					

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EXAMPLE 6 Determining the chirality configuration of astaxanthin produced in heterologous systems

The chirality configurations of astaxanthin produced by *Escherichia coli* cells, as described under Example 3 hereinabove, and by cyanobacterium *Synechococcus* PCC7942 cells, as described in Example 5 hereinabove, were determined by HPLC of the derived diastereoisomeric camphanates of the astaxanthin [Renstrom B, Borch G, Skulberg M and Liaaen-Jensen S (1981) Optical purity of (3S,3S')-astaxanthin from *Haematococcus pluvialis*. Phytochem 20: 2561-2565]. The analysis proved that the *Escherichia coli* and *Synechococcus* PCC7942 cells described above, synthesize pure (3S,3'S) astaxanthin.

EXAMPLE 7 Transformation of a higher plant with crtO

Producing natural astaxanthin in higher plants has two anticipated benefits. First, as a pure chemical, astaxanthin is widely used as feed additive for fish. It is a potential food colorant suitable for humans consumption and has potential applications in the cosmetic industry. Second, inducing astaxanthin biosynthesis *in vivo* in flowers and fruits will provide attractive pink/red colors which will increase their appearance and/or nutritious worth.

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In flowers and fruits carotenoids are normally synthesized and accumulated to high concentration in chromoplasts, a typical pigment-containing plastids, thus providing typical intense colors to these organs. Inducing synthesis of astaxanthin in chromoplasts enables the accumulation of high concentration of this ketocarotenoid. Over-expression of carotenoid biosynthesis genes which results in elevated concentrations of carotenoids in chloroplasts, or other alterations in carotenoid composition in chloroplasts may damage the thylakoid membranes, impair photosynthesis and thus is deleterious to the plants. In contrast, increase of carotenoid concentration or alteration in carotenoid composition in chromoplasts do not affect the viability of the plant nor the yield of fruits and flowers.

Thus, gene-transfer technology was used to implant the *crtO* gene isolated from the alga *Haematococcus pluvialis*, as described, into a higher plant, in such a way that its expression is up-regulated especially in chromoplast-containing cells.

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To this end, a T-DNA containing binary plasmid vector as shown in Figure 12 was assembled in *E. coli* from the promoter and coding DNA sequences of the transit peptide encoded by the *Pds* gene from a tomato species *Lycopersicon esculentum*, linked to the coding DNA sequence of *crtO* from *H. pluvialis*. Upon stable transfer of this DNA construct via *Agrobacterium*-mediated transformation into a tobacco (*Nicotiana tabacum NN*) plant to form a transgenic plant, as described under methods above, the plant acquired the ability to produce ketocarotenoids especially in flower tissues (chromoplast-containing cells). It should be noted that the *Pds* gene promoter is capable of directing transcription and therefore expression especially in chloroplasts and/or chromoplasts-containing tissues of plants. It should be further noted that the transit peptide encoded by part of the *Pds* coding sequence is capable of directing conjugated (i.e., in frame) proteins into plant chromoplasts and/or chloroplasts.

As shown in Figure 15, in chromoplasts-containing cells, such as in the nectary tissue of the flower of tobacco, this DNA construct induces accumulation of astaxanthin and other ketocarotenoids to a higher level which alters the color from the normal yellow to red.

Concentration and composition of carotenoids in chloroplasts-containing tissues, such as leaves, and in chromoplast-containing tissues, such as flowers, were determined in the transgenic plants and compared to normal non-transformed plants.

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Carotenoids compositions in leaves (chloroplasts-containing tissue) and in the nectary tissue of flowers (chromoplast containing tissue) of wild type and transgenic tobacco plants were determined by thin layer chromatography (TLC) and by high pressure liquid chromatography (HPLC) as described above.

Total carotenoids concentration in leaves (chloroplasts-containing tissue) and in the nectary tissue of flowers (chromoplast containing tissue) of wild type and transgenic tobacco plants are summarized in Tables 3 below.

Percents of carotenoids composition in leaves of wild-type and transgenic tobacco plants are summarized in Tables 4 below.

Percents of carotenoids composition in the nectary tissue of flowers of wild-type and transgenic tobacco plants are summarized in Tables 5 below.

TABLE 3

µg carotenoids per gr fresh weight

		Wild-type	Transgenic with crtO
20	Leaf (Chloroplasts)	200	240
	Nectary tissue (Chromoplasts)	280	360

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TABLE 4
% of total carotenoids composition in chloroplasts-containing tissue (leaf)
Wild-type Transgenic

β-carotene	29.9	26.7
neoxanthin	5.0	5.9
violaxanthin	11.6	18.1
antheraxanthin	4.9	2.6
lutein	43.9	41.4
zeaxanthin	4.7	4.3
astaxanthin + adonirubin	0.0	1.0

TABLE 5
% of total carotenoid composition in chromoplasts-containing tissue (flower)

		Wild-type	Transgenic
5			
	beta-carotene	58.1	21.0
	violaxanthin	40.3	1.5
	lutein	0.0	1.1
	zeaxanthin	1.6	1.0
10	hydroxyechinenone	0.0	13.7
	3'hydroxyechinenone	0.0	4.1
	adonirubin	0.0	22.4
	adonixanthin	0.0	8.7
	astaxanthin	0.0	26.5

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Please note the elevated content of hydroxyechinenone, 3'hydroxyechinenone, adonirubin, adonixanthin and astaxanthin especially in the chromoplast containing tissue of the transgenic tobacco plants.

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Thus, the present invention successfully addresses the shortcomings of the presently known configurations by enabling a relatively low cost biotechnological production of (3S,3'S) astaxanthin by providing a peptide having a β -C-4-oxygenase activity; a DNA segment coding for this peptide; an RNA segments coding for this peptide; a recombinant DNA molecule comprising a vector and the DNA segment; a host containing the above described recombinant DNA molecule or DNA segment; and of a method for biotechnologically producing (3S,3'S) astaxanthin or a food additive containing (3S,3'S) astaxanthin, using the host.

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While the invention has been described with respect to a limited number of embodiments, it will be appreciated that many variations, modifications and other applications of the invention may be made.

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SEQUENCE LISTING

			SEQUENCE LISTING
(1)	GENERAL	INFORMATION:	
	(i)	APPLICANTS:	Joseph Hirschberg, Tamar Lotan and
			Mark Harker
	(ii)	TITLE OF INVENTION:	Polynucleotide molecule from
			Haematococcus pluvialis encoding a polypeptide having a β -C-4-oxygenase
			activity for biotechnological production of
			(3S,3'S) astaxanthin.
	(iii)	NUMBER OF SEQUENCES:	4
	(iv)	CORRESPONDENCE ADDRESS:	
		(A) ADDRESSEE:	Mark M. Friedman c/o Robert Sheinbein
		(B) STREET:	2940 Birchtree space lane
		(C) CITY:	Silver Spring
		(D) STATE:	Maryland
		(E) COUNTRY:	United States of America
		(F) ZIP:	20906
	(v)	COMPUTER READABLE FORM:	
		(A) MEDIUM TYPE:	1.44 megabyte, 3.5" microdisk
		(B) COMPUTER:	Twinhead Slimnote-890TX
		(C) OPERATING SYSTE	•
			Windows version 3.11
		(D) SOFTWARE:	Word for Windows version 2.0
	(vi)	CURRENT APPLICATION DATA	
		(A) APPLICATION NUM	BEK:
		(B) FILING DATE:	
	(viii)	(C) CLASSIFICATION: PRIOR APPLICATION DATA:	
	(vii)	(A) APPLICATION NUM	RED.
		(B) FILING DATE:	DEN.
	(viii)	ATTORNEY/AGENT INFORMAT	ION:
	,	(A) NAME:	Friedmam, Mark M.
		(B) REGISTRATION NU	`
		(C) REFERENCE/DOCKE	·
	(ix)	TELECOMMUNICATION INFOR	MATION:
		(A) TELEPHONE:	972-3-5625553
		(B) TELEFAX:	972-3-5625554
		(C) TELEX:	
(2)	INFORMA	ATION FOR SEQ ID NO:1:	
	(i)	SEQUENCE CHARACTERISTIC	S:
		(A) LENGTH:	1771 base pairs
		(B) TYPE:	nucleic acid
		(C) STRANDEDNESS:	double
		(D) TOPOLOGY:	linear
	(xi)	SEQUENCE DESCRIPTION:	SEQ ID NO:1:
GGC A	CG AGC TI	TG CAC GCA AGT CAG CGC GC	G CAA GTC AAC ACC TGC CGG 48
TCC A	CA GCC TO	CA AAT AAT AAA GAG CTC AA	G CGT TTG TGC GCC TCG ACG 96
TGG C	CA GTC TO	GC ACT GCC TTG AAC CCG CG	A GTC TCC CGC CGC ACT GAC 144
TGC C	AT AGC AG	CA GCT AGA CGA ATG CAG CT.	A GCA GCG ACA GTA ATG TTG 192
GAG C	AG CTT AG	CC GGA AGC GCT GAG GCA CT	C AAG GAG AAG GAG 240
GTT G	CA GGC A	GC TCT GAC GTG TTG CGT AC	A TGG GCG ACC CAG TAC TCG 288
			C CCG GGA CTG AAG AAT GCC 336
			C ATC ACA ATG GCG CTA CGT 384
			C CAC GCC ATT TTT CAA ATC 432
			C TGG CTG CCC GTG TCA GAT 480
GCC A	CA GCT C	AG CTG GTT AGC GGC ACG AG	C AGC CTG CTC GAC ATC GTC 528

												•				
GTA	GTA	TTC	TTT	GTC	CTG	GAG	TTC	CTG	TAC	ACA	GGC	CTT	TTT	ATC	ACC	576
ACG	CAT	GAT	GCT	ATG	CAT	GGC	ACC	ATC	GCC	ATG	AGA	AAC	AGG	CAG	CTT	624
AAT	GAC	TTC	TTG	GGC	AGA	GTA	TGC	ATC	TCC	TTG	TAC	GCC	TGG	TTT	GAT	672
TAC	AAC	ATG	CTG	CAC	CGC	AAG	CAT	TGG	GAG	CAC	CAC	AAC	CAC	ACT	GGC	720
GAG	GTG	GGC	AAG	GAC	CCT	GAC	TTC	CAC	AGG	GGA	AAC	CCT	GGC	ATT	GTG	768
CCC	TGG	TTT	GCC	AGC	TTC	ATG	TCC	AGC	TAC	ATG	TCG	ATG	TGG	CAG	TTT	816
GCG	CGC	CTC	GCA	TGG	TGG	ACG	GTG	GTC	ATG	CAG	CTG	CTG	GGT	GCG	CCA	864
ATG	GCG	AAC	CTG	CTG	GTG	TTC	ATG	GCG	GCC	GCG	CCC	ATC	CTG	TCC	GCC	912
TTC	CGC	TTG	TTC	TAC	TTT	GGC	ACG	TAC	ATG	CCC	CAC	AAG	CCT	GAG	CCT	960
GGC	GCC	GCG	TCA	GGC	TCT	TCA	CCA	GCC	GTC	ATG	AAC	TGG	TGG	AAG	TCG	1008
CGC	ACT	AGC	CAG	GCG	TCC	GAC	CTG	GTC	AGC	TTT	CTG	ACC	TGC	TAC	CAC	1056
TTC	GAC	CTG	CAC	TGG	GAG	CAC	CAC	CGC	TGG	CCC	TTC	GCC	CCC	TGG	TGG	1104
GAG	CTG	CCC	AAC	TGC	CGC	CGC	CTG	TCT	GGC	CGA	GGT	CTG	GTT	CCT	GCC	1152
TAG	CTG	GAC	ACA	CTG	CAG	TGG	GCC	CTG	CTG	CCA	GCT	GGG	CAT	GCA	GGT	1200
TGT,	GGC	AGG	ACT	GGG	TGA	GGT	GAA	AAG	CTG	CAG	GCG	CTG	CTG	CCG	GAC	1248
ACG	CTG	CAT	GGG	CTA	CCC	TGT	GTA	GCT	GCC	GCC	ACT	AGG	GGA	GGG	GGT	1296
TTG	TAG	CTG	TCG	AGC	TTG	CCC	CAT	GGA	TGA	AGC	TGT	GTA	GTG	GTG	CAG	1344
GGA	GTA	CAC	CCA	CAG	GCC	AAC	ACC	CTT	GCA	GGA	GAT	GTC	TTG	CGT	CGG	1392
GAG	GAG	TGT	TGG	GCA	GTG	TAG	ATG	CTA	TGA	TTG	TAT	CTT	AAT	GCT	GAA	1440
GCC	TTT	AGG	GGA	GCG	ACA	CTT	AGT	GCT	GGG	CAG	GCA	ACG	CCC	TGC	AAG	1488
GTG	CAG	GCA	CAA	GCT	AGG	CTG	GAC	GAG	GAC	TCG	GTG	GCA	GGC	AGG	TGA	1536
AGA	GGT	GCG	GGA	GGG	TGG	TGC	CAC	ACC	CAC	TGG	GCA	AGA	CCA	TGC	TGC	1584
AAT	GCT	GGC	GGT	GTG	GCA	GTG	AGA	GCT	GCG	TGA	TTA	ACT	GGG	CTA	TGG	1632
ATT	GTT	TGA	GCA	GTC	TCA	CTT	ATT	CTT	TGA	TAT	AGA	TAC	TGG	TCA	GGC	1680
						TGA					GTG	GTG	CGC	TGC	CCC	1728
TGC	GCT	TAT	GAA	GCT	GTA	ACA	ATA	AAG	TGG	TTC						1771

(2) INFORMATION FOR SEQ ID NO:2:

(xi)

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1771 bases
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear
SEQUENCE DESCRIPTION: SEQ ID NO:2:

GGC ACG AGC UUG CAC GCA AGU CAG CGC GCG CAA GUC AAC ACC UGC CGG 48 UCC ACA GCC UCA AAU AAU AAA GAG CUC AAG CGU UUG UGC GCC UCG ACG 96 UGG CCA GUC UGC ACU GCC UUG AAC CCG CGA GUC UCC CGC CGC ACU GAC 144 UGC CAU AGC ACA GCU AGA CGA AUG CAG CUA GCA GCG ACA GUA AUG UUG 192 GAG CAG CUU ACC GGA AGC GCU GAG GCA CUC AAG GAG AAG GAG AAG GAG 240 GUU GCA GGC AGC UCU GAC GUG UUG CGU ACA UGG GCG ACC CAG UAC UCG 288 CUU CCG UCA GAA GAG UCA GAC GCG GCC CGC CCG GGA CUG AAG AAU GCC UAC AAG CCA CCA CCU UCC GAC ACA AAG GGC AUC ACA AUG GCG CUA CGU GUC AUC GGC UCC UGG GCC GCA GUG UUC CUC CAC GCC AUU UUU CAA AUC 432 AAG CUU CCG ACC UCC UUG GAC CAG CUG CAC UGG CUG CCC GUG UCA GAU 480 GCC ACA GCU CAG CUG GUU AGC GGC ACG AGC AGC CUG CUC GAC AUC GUC 528 GUA GUA UUC UUU GUC CUG GAG UUC CUG UAC ACA GGC CUU UUU AUC ACC 576 ACG CAU GAU GCU AUG CAU GGC ACC AUC GCC AUG AGA AAC AGG CAG CUU 624 AAU GAC UUC UUG GGC AGA GUA UGC AUC UCC UUG UAC GCC UGG UUU GAU 672 UAC AAC AUG CUG CAC CGC AAG CAU UGG GAG CAC CAC AAC CAC ACU GGC 720 GAG GUG GGC AAG GAC CCU GAC UUC CAC AGG GGA AAC CCU GGC AUU GUG 768 CCC UGG UUU GCC AGC UUC AUG UCC AGC UAC AUG UCG AUG UGG CAG UUU GCG CGC CUC GCA UGG UGG ACG GUG GUC AUG CAG CUG CUG GGU GCG CCA 864 AUG GCG AAC CUG CUG GUG UUC AUG GCG GCC GCG CCC AUC CUG UCC GCC 912 UUC CGC UUG UUC UAC UUU GGC ACG UAC AUG CCC CAC AAG CCU GAG CCU 960 GGC GCC GCG UCA GGC UCU UCA CCA GCC GUC AUG AAC UGG UGG AAG UCG 1008 CGC ACU AGC CAG GCG UCC GAC CUG GUC AGC UUU CUG ACC UGC UAC CAC 1056

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											44	ļ				
uuc d	GAC	CUG	CAC	UGG	GAG (CAC	CAC (CGC (JGG (CCC	uuc	GCC	CCC I	UGG	UGG	1104
GAG (CUG	CCC	AAC	UGC	CGC (CGC (ו פעב	JCU (GGC	CGA	GGU	CUG	GUU (CCU	GCC	1152
UAG (CUG	GAC	ACA	CUG	CAG (JGG (CC (cug (CUG	CCA	GCU	GGG	CAU	GCA	GGU	1200
UGU (GGC	AGG	ACU	GGG	UGA (GGU (GAA A	AAG (CUG	CAG	GCG	CUG	CUG	CCG	GAC	1248
ACG (CUG	CAU	GGG	CUA	CCC I	JGU (GUA (GCU	GCC	GCC	ACU	AGG	GGA	GGG	GGU	1296
UUG (1344
					GCC											1392
	_				GUG (1440
					ACA											1488
					AGG											1536
-					UGG I											1584
					GCA											1632
					UCA											1680
					GUA						GUG	GUG	CGC	UGC	CCC	1728
UGC	GCU	UAU	GAA	GCU	GUA	ACA .	AUA	AAG	UGG	UUC						1771
							- N	7 .								
(2)					FOR S				T 1 CC	_						
		(i)			ENCE			CKIS	1102		'1 br		airs			
				(A)		LENG TYPE						se p				
				(B)		STRA		MESS			ıble		u			
				(D)		TOPO			••	doc		Li	near			
		(xi)			JENCE				•	SEC) ID	NO:3				
		(JLWC	LNOL	020			•	-			•			
GGC	ACG	AGC	TTG	CAC	GCA	AGT	CAG	CGC	GCG	CAA	GTC	AAC	ACC	TGC	CGG	48
					AAT											96
					GCC											144
					AGA											192
													Val			
											5					
GAG	CAG	CTT	ACC	GGA	AGC	GCT	GAG	GCA	CTC	AAG	GAG	AAG	GAG	AAG	GAG	240
Glu	Gln	Leu	Thr	Gly	Ser	Ala	Glu	Ala	Leu	Lys	Glu	Lys	Glu	Lys	Glu	
10					15					20					25	
					GAC											288
Val	Ala	Gly	Ser	Ser	Asp	Val	Leu	Arg	Thr	ĭrp	Ala	Thr	Gln	Tyr	Ser	
				30					35					40		
					TCA											336
Leu	Pro	Ser	Glu	Glu	Ser	Asp	Ala			Pro	Gly	/ Leu			Ala	
			45					50					55			
					TCC											384
Tyr	Lys			Pro	Ser	Asp			Gly	116	ומו			rec	Arg	
		60					65					70		CAA	ATC	432
					GCC											436
val			/ Ser	. itt	Ala			Pne	Leu	пі	8!		Pile		1 116	
440	75			. TC	C TTG	600		CTC	. רמר	TG			CTO	: TC	CAT	480
															Asp	400
90 90		J Pire	, ,,,,	361	95			LCC		100			, ,,,,		105	
		7 GC.	T (^A)	; CT:			GGC	: ACG	: AGC			G CT	C GAC	: AT	GTC	528
															e Val	
A 1 6	,,,			110			٠.,		115					12		
GTA	GTA	A TT	с тт			GAG	TT0	: ст			A GG	с ст	T T T		CACC	576
															e Thr	
			12!					130					135			
ACC	CA	T GA			G CA1	GGC	: AC	CATO	C GC	CAT	G AG	A AA	C AG	G CA	G CTT	624
															n Leu	
		14					14!					15				

140 145 150

AAT	GAC	TTC	TTG	GGC	AGA	GTA	TGC	ATC	TCC	TTG	TAC	GCC	TGG	TTT	GAT	672
Asn	Asp	Phe	Leu	Gly	Arg	Val	Cys	Ιle	Ser	Leu	Tyr	Ala	Trp	Phe	Asp	
	155					160					165					
TAC	AAC	ATG	CTG	CAC	CGC	AAG	CAT	TGG	GAG	CAC	CAC	AAC	CAC	ACT	GGC	720
Tyr	Asn	Met	Leu	His	Arg	Lys	His	Trp	Glu	His	Kis	Asn	His	Thr	Gly	
170					175					180					185	
GAG	GTG	GGC	AAG	GAC	CCT	GAC	TTC	CAC	AGG	GGA	AAC	CCT	GGC	ATT	GTG	768
Glu	Val	Gly	Lys	Asp	Pro	Asp	Phe	His	Arg	Gly	Asn	Pro	Gly	Ile	Val	
				190					195					200		
CCC	TGG	TTT	GCC	AGC	TTC	ATG	TCC	AGC	TAC	ATG	TCG	ATG	TGG	CAG	TTT	816
Pro	Trp	Phe	Ala	Ser	Phe	Met	Ser	Ser	Туг	Met	Ser	Met	Тгр	Gln	Phe	
			205					210					215			
GCG	CGC	CTC	GCA	TGG	TGG	ACG	GTG	GTC	ATG	CAG	CTG	CTG	GGT	GCG	CCA	864
Ala	Arg	Leu	Ala	Тгр	Тгр	Thr	Val	Val	Met	Gln	Leu	Leu	Gly	Ala	Pro	
		220					225					230				
		AAC														912
Met	Ala	Asn	Leu	Leu	Val	Phe	Met	Ala	Ala	Ala	Pro	Ile	Leu	Ser	Ala	
	235					240					245					
TTC	CGC	TTG	TTC	TAC	TTT	GGC	ACG	TAC	ATG	CCC	CAC	AAG	CCT	GAG	CCT	960
Phe	Arg	Leu	Phe	Туг	Phe	Gly	Thr	Tyr	Met	Pro	His	Lys	Pro	Glu	Pro	
250					255					260					265	
		GCG														1008
Gly	Ala	Ala	Ser		Ser	Ser	Pro	Ala	Val	Met	Asn	Trp	1 rp	Lys	Ser	
				270					275					280		
		AGC														1056
Arg	Thr	Ser		Ala	Ser	Asp	Leu		Ser	Phe	Leu	Thr		Tyr	His	
			285					290					295			
		CTG														1104
Phe	Asp	Leu	His	Trp	Glu	His		Arg	Trp	Pro	Phe		Pro	Ĭгр	Trp	
		300					305					310				
		CCC														1152
GLU		Pro	Asn	Cys	Arg	_	Leu	Ser	Gly	Arg		Leu	Val	Pro	Ala	
	315					320					325					
		GAC														1200
		AGG														1248
		CAT														1296
		CTG														1344
		CAC														1392
		TGT														1440
		AGG														1488
		GCA														1536
		GCG														1584
		GGC														1632
		TGA														1680
		GGA									GIG	GIG	LGC	IGC	CCC	1728
IUC	GC [TAT	GAA	GCT	GIA	ACA	AIA	AAG	1 GG	110						1771

(2) INFORMATION FOR SEQ ID NO:4:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 329 amino acids
(B) TYPE: amino acid

(B) TYPE: amino aci

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Gin Leu Ala Ala Thr Val Met Leu

5

Glu Gln Leu Thr Gly Ser Ala Glu Ala Leu Lys Glu Lys Glu Lys Glu

10					15					20					25
Val	Ala	Gly	Ser	Ser 30	Asp	Val	Leu	Arg	Thr 35	Trp	Ala	Thr	Gln	Tyr 40	Ser
Leu	Pro	Ser	Glu 45	Glu	Ser	Asp	Ala	Ala 50	Arg	Pro	Gly	Leu	Lys . 55	Asn	Ala
Tyr	Lys	Pro 60	Pro	Pro	Ser	Asp	Thr 65	Lys	Gly	Ile	Thr	Met 70	Ala	Leu	Arg
Val	11e 75	Gly	Ser	Trp	Ala	Ala 80	Val	Phe	Leu	His	Ala 85	Ile	Phe	Gln	Ile
Lys 90	Leu	Pro	Thr	Ser	Leu 95	Asp	Gln	Leu	His	Тгр 100	Leu	Pro	Val	Ser	Asp 105
				110					115				Asp	120	
Val	Val	Phe	Phe 125	Val	Leu	Glu	Phe	Leu 130	Tyr	Thr	Gly	Leu	Phe 13 5	Ile	Thr
Thr	His	Asp 140	Ala	Met	His	Gly	Thr 145	Ile	Ala	Met	Arg	Asn 150	Arg	Gln	Leu
Asn	Asp 155	Phe	Leu	Gly	Arg	Val 160	Cys	Ile	Ser	Leu	Tyr 165	Ala	Trp	Phe	Asp
170					175	·				180			His		185
Glu	Val	Gly	Lys	Asp 190	Pro	Asp	Phe	His	Arg 195	Gly	Asn	Pro	Gly	1 l e 200	Val
Pro	Trp	Phe	Ala 205	Ser	Phe	Met	Ser	Ser 210	Туг	Met	Ser	Met	Trp 215	Gln	Phe
Ala	Arg	Leu 220	Ala	Тгр	Тгр	Thr	Val 225	Val	Met	Gln	Leu	Leu 230	Gly	Ala	Pro
Met	Ala 235	Asn	Leu	Leu	Val	Phe 240	Met	Ala	Ala	Ala	Pro 245	Ile	Leu	Ser	Ala
Phe 250	Arg	Leu	Phe	Туг	Phe 255	Gly	Thr	Туг	Met	Pro 260	His	Lys	Pro	Glu	Pro 265
Gly	Ala	Ala	Ser	Gly 270	Ser	Ser	Рго	Ala	Va l 275	Met	Asn	Тгр	Тгр	Lys 280	Ser
Arg	Thr	Ser	Gln 285	Ala	Ser	Asp	Leu	Val 290	Ser	Phe	Leu	Thr	Cys 295	Туг	His
Phe	Asp	Leu 300		Тгр	Glu	His	His 305	Arg	Тrр	Pro	Phe	Ala 310	Pro	Trp	Trp
Glu	Leu 315	Pro	Asn	Cys	Arg	Arg 320		Ser	Gly	Arg	Gly 325	Leu	Val	Pro	Ala

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WHAT IS CLAIMED IS:

- 1. A DNA segment comprising a nucleotide sequence coding for a polypeptide having an amino acid sequence as set forth in SEQ ID NO:4
- 2. A DNA segment as in claim 1, wherein said nucleotide sequence is a variant selected from the group of variants consisting of allelic variants, species variants, naturally occurring variants, man-induced variants and combinations thereof.
- 3. A DNA segment as in claim 1, wherein said nucleotide sequence includes a sequence as set forth in SEQ ID NO:1.
- 4. A DNA segment as in claim 1, wherein said nucleotide sequence includes a sequence as set forth between and including nucleotides 166 and 1152 of SEQ ID NO:1.
- 5. A DNA segment comprising a nucleotide sequence hybridizing under high stringency conditions to a nucleic acid probe, said probe including at least fifteen successive nucleotides of SEQ ID NO:1.
- 6. A DNA segment comprising a nucleotide sequence hybridizing under low-stringency conditions to a nucleic acid probe, said probe including nucleotides 166 through 1152 of SEQ ID NO:1.
- 7. A DNA segment comprising a nucleotide sequence hybridizing under high stringency conditions to a nucleic acid probe, said probe including at least fifteen successive nucleotides of SEQ ID NO:2.
- 8. A DNA segment comprising a nucleotide sequence hybridizing under low-stringency conditions to a nucleic acid probe, said probe including nucleotides 166 through 1152 of SEQ ID NO:2.
- 9. An RNA segment comprising a nucleotide sequence coding for a polypeptide having an amino acid sequence as set forth in SEO ID NO:4
- 10. An RNA segment as in claim 9, wherein said nucleotide sequence is a variant selected from the group of variants consisting of allelic variants, species

variants, naturally occurring variants, man-induced variants and combinations thereof.

- 11. An RNA segment as in claim 9, wherein said nucleotide sequence includes a sequence as set forth in SEQ ID NO:2.
- 12. An RNA segment as in claim 9, wherein said nucleotide sequence includes a sequence as set forth between and including nucleotides 166 and 1152 of SEQ ID NO:2.
- 13. An RNA segment comprising a nucleotide sequence hybridizing under high stringency conditions to a nucleic acid probe, said probe including at least fifteen successive nucleotides of SEQ ID NO:1.
- 14. An RNA segment comprising a nucleotide sequence hybridizing under low-stringency conditions to a nucleic acid probe, said probe including nucleotides 166 through 1152 of SEQ ID NO:1.
- 15. An RNA segment comprising a nucleotide sequence hybridizing under high stringency conditions to a nucleic acid probe, said probe including at least fifteen successive nucleotides of SEQ ID NO:2.
- 16. An RNA segment comprising a nucleotide sequence hybridizing under low-stringency conditions to a nucleic acid probe, said probe including nucleotides 166 through 1152 of SEQ ID NO:2.
- 17. A polypeptide comprising an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene, allelic and species variants, and functional naturally occurring and man-induced variants thereof.
- 18. A polypeptide as in claim 17, wherein said amino acid sequence is as set forth in SEQ ID NO:4.
- 19. A polypeptide comprising an amino acid sequence homologous to the sequence set forth in SEQ ID NO:4.
- 20. A polypeptide comprising an amino acid sequence being encoded by a DNA segment, said DNA segment hybridizing under low stringency conditions

to nucleotides 166 through 1152 of SEQ ID NO:1, the polypeptide having a β -C-4-oxygenase activity.

- 21. A recombinant vector DNA molecule comprising a DNA segment as in claim 2.
- 22. A host comprising a recombinant vector DNA molecule as in claim 21, said host is selected from the group consisting of a cell and an organism.
- 23. A host comprising a DNA segment as in claim 2, said host is selected from the group consisting of a cell and an organism.
- 24. A method of producing xanthophylls selected from the group consisting of astaxanthin, canthaxanthin, echinenone, cryptoxanthin, isocryptoxanthin, hydroxyechinenone, zeaxanthin, adonirubin or adonixanthin and combinations thereof, comprising the steps of:
 - (a) providing a host as in claim 22;
 - (b) providing said host with growing conditions for production of the xanthophylls; and
 - (c) extracting the xanthophylls from said host.
- 25. A method of producing xanthophylls selected from the group consisting of astaxanthin, canthaxanthin, echinenone, isocryptoxanthin, cryptoxanthin, hydroxyechinenone, zeaxanthin, adonirubin or adonixanthin and combinations thereof, comprising the steps of:
 - (a) providing a host as in claim 23;
 - (b) providing said host with growing conditions for production of the xanthophylls; and
 - (c) extracting the xanthophylls from said host.
 - 26. A host as in claim 22, wherein said host is used as a food additive.
 - 27. A host as in claim 23, wherein said host is used as a food additive.
- 28. A transgenic plant expressing a transgene coding for a polypeptide including an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene, allelic and species variants or functional naturally occurring or man-induced variants thereof.

- 29. A transgenic plant as in claim 28, wherein said expression is highest in chromoplasts-containing tissues.
- 30. A recombinant DNA vector comprising a first DNA segment encoding a polypeptide for directing a protein into plant chloroplasts or chromoplasts and an in frame second DNA segment encoding a polypeptide including an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene, allelic and species variants or functional naturally occurring and maninduced variants thereof.
- 31. A recombinant DNA vector as in claim 30, wherein said first DNA segment is derived from the *Pds* gene of tomato.
- 32. A recombinant DNA vector comprising a first DNA segment including a promoter highly expressible in plant chloroplasts or chromoplasts-containing tissues and a second DNA segment encoding a polypeptide including an amino acid sequence corresponding to *Haematococcus pluvialis crtO* gene, allelic and species variants or functional naturally occurring and man-induced variants thereof.
- 33. A recombinant DNA vector as in claim 30, wherein said first DNA segment is derived from the *Pds* gene of tomato.

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FIG. 2-A

					_	
CRTOA.SEQ	ACCACCTTACCCC	210 MCCCCTCAC	220	CYCYYCCYC-Y	240 UCCACCTTC	250 CACCCACC
CRTOJ.SEQ	ATCCACCTCCCAT	CCCCCCTAAT	CCTCCACCA		CACCCACCTCC	TTCCXCC 60
CRTOA.SEQ	260 TCTCACCTCTTCC	20 270 ·	280 280	290	50 300 TACAACACTI	310
CKIOK.3CQ	: ::::: :::					
CRTOJ.SEQ	CCAGACGTCTTGA 70	80	90	100	110	120
CRTOA.SEQ	CCCCCCCCCCAC					
CRTOJ.SEQ	CCTCCTCCTCCCCC					
	380	390	400	410	420	430
CRTOA.SEQ	ATCCCCCTACCTC					
CXTOJ.SEQ	ATCCCCCTCACCA			220		
	440	450	460	470	480	490
CRTOA.SEQ	ACCTTCCCACCT					
CRTOJ.SEQ	ACCCTACCGACAT					
••	250	260	270	280	290	300
CRTOA.SEQ	SOO CTCCTTACCCCCA	210 הכניאכראניבר	520 IGCTUGACAT	530 CGTCGTAGTA	540	550 TGGAGTTG
CKIOK.JEQ			_	:: ::::		
CRT01.SEQ	310	120 320	TACTCCACAT 330	CCCTCCACTC 340	TCATTCTAC 350	360
	560	570	580	590	600	610
CRTOA.SEQ	CTCTACACACCCC					
CRTOJ.SEQ	CTCTACACTCCTC					
	370 620	380 630	390 640	400 650	410 660	420 670
CRTOA.SEQ	MCACCCACCTTA	_				
Comi SEO	:::::::::::::::::::::::::::::::::::::::					
CRTOJ.SEQ	CACACCCACCTC 430	440	450	460	470	480
	680	690	700	710	720	730
CRTOA.SEQ	TACAACATCCTCC					
CRTOJ.SEQ	TACACCATCCTCC	CATCCCLACCA	ACTOCCACCA	CCACAACCAT	ACTECEGAAC	TCCCCALLA
	490 740	5C0 750	510 760	520 770	530 780	540 790
CRTOA.SEQ	CACCCTGACTTCC					
CATOJ.SEQ	GACCCTGACTTC					
CK103.3EQ	550	560	570	580	590	600
CD704 570	800	610		830	840	850
CRTOA.SEQ	ACCTACATGTCG	-		CCCXTCCTCC		
D3S.LOTKO	ACCTACATGTCCC	TOTOGOLAGT	TCCCCCCCT	CCCATCCTCC	CCACTCCTCA	TCCLUSTC
	610 860	629 870	630 830	640 830	650 900	660 910
CRTOA.SEQ	CTCCCTG-CCCA					-
CATOJ.SEO	CTCCCCCCCCC.		_			
CR107.364	670	680	690	700	710	720
C2704 550	920	930	940	950	960	970
CXTOA.SEQ	TTCCCCTTCTTC					
D32.1CTKO	730					

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FIG. 2-B

	980	990	1000	1010	1020	1030
CRTOA.SEQ	CCCTCTTCACCA	CCCCTCATCA	MCTCCTCCA	ACTCCCCAC	TACCCACCCS.	TCCCACCTC
	::::: ::	: :: :::	::::: :	:: ::	11 11	
CSTOJ.SEQ	CCCTCTCA	CCTCATC(CCTCCTTCA	CCCCAACAC	MCTCACCCA	TCTCATGTG
		790	800	810	820	830
	1040	1050	1060	1070	1080	1090
CRTOA.SEQ	CTCACCTTTCTC	ACCTCCTAC	CACTTCGACC	TCCACTCCCA	CCACCACCCC	TCCCCCTTC
	: :: :: :::	:: ::::::		:::::::::	::::::: :	:::::::
D32. LOTKO	ATGAGTTTCCTC	CATCCTAC	CACTTTGACC	TCCACTCCCA	CCACCACACC.	TCCCCCTTT
	840	850	860	870	880	890
	1100	1110	1120			
CRTOA.SEQ		1110	1120	1130	1140	1150
C. 107.31.4	CCCCCTCCTCC	~~~	MCIUCUC	accrete tete	CCCACCTCTC	CTTCCTCCC
CATOJ.SEQ	CCCCCTCCTC		-::::::::	<u> </u>	:: :: :::	:: ::::::
G	900	910				
	1160	1170	920	930	940	950
CRTOA.SEQ			1130	119	0 120	0
	TACCTCCACACA	:::				
CX.LO1.2EO	TTGCCATCAC		: ::: ::	::::::	::: ::	:::
	960	970	980			
	1210 1220		760	990	1000	1010
CRTOA.SEQ	ACTECETEACET					
	:: :::::	•				
CRTOJ.SEQ	ACACCCTCCTCA	ı				
	1020	•				
	1010					

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FIG. 3

CRTOA.AMI CRTOJ.AMI	: :	: :: ::	::	: : ::::	:::: :::	50 ESDAARPGLKNAY ::::::::::::::::::::::::::::::::::::
CRTOA.AMI	60 KPPPSD	70 TKGITHALR	80 VIGSVAAVFL	90 HAIFOIKLPT:	100 SLDOLKWLPV:	110 SDATAQLVSGTSSLL
0	::: ::	::::::	:: : ::::	::::::	: :::::::::	: ::::: : ::::
CRTOJ.AHI	KPPASD 50	AKGITHALT 60	TIGTWTAVFL 70	HAIFQIRLPT 80	SHDQLHWLPV: 90	SEATAQLLGGSSSLL 100
	120	130	140	150	160	170
CRTOA.AMI				GTIAHRNRQL :::: : :::		LYAVFDYMLHRKHW ::::::::
CRTOJ.AMI						LYAWFDYSHLHRKHW 160
	180	190	200	210	220	230
CRTOA.AMI						TVVHQLLGAPKANLL
CRTOJ.AHI	170	180	190	200	210	AVVHQHLGAPHANLL 220
	240	250	260	270	280	290
CRTOA.AMI						SQASDLVSFLT
CRTOJ.AMI			ETTECTYLPHKE Z50		:: : : : VMAWFRAKT 270	SEASDVMSFLT 280
CRTOA.AMI	300 Cyhfdi	LHWEHHRWP	FAPWWELPHC	20 329 RRLSGRGLVP4	X	
CRTOJ.AMI		LHWEHHRWP	FAPHHQLPHC	RRLSGRGLVPA		

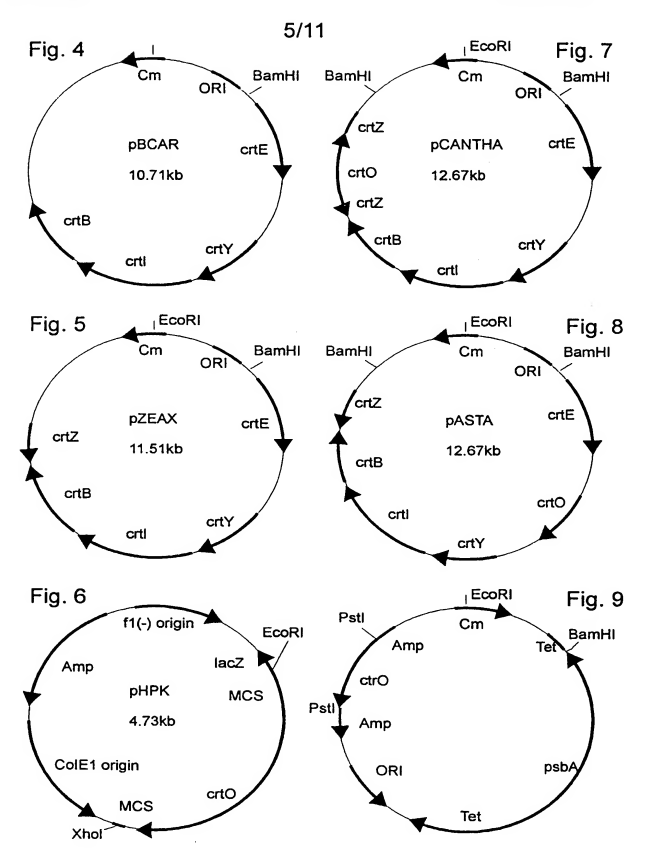


Fig. 10

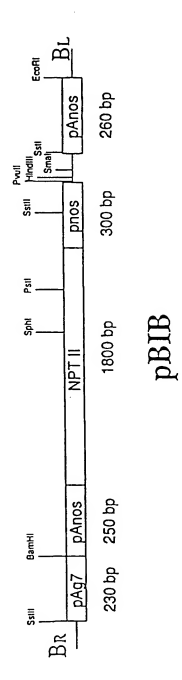


Fig. 11

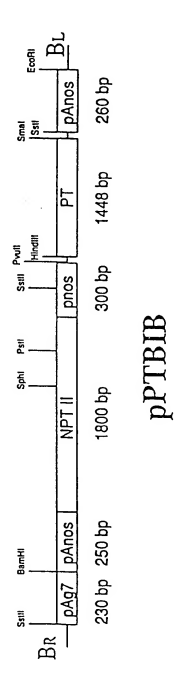


Fig. 12

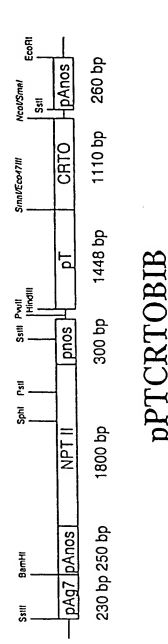


Figure 13

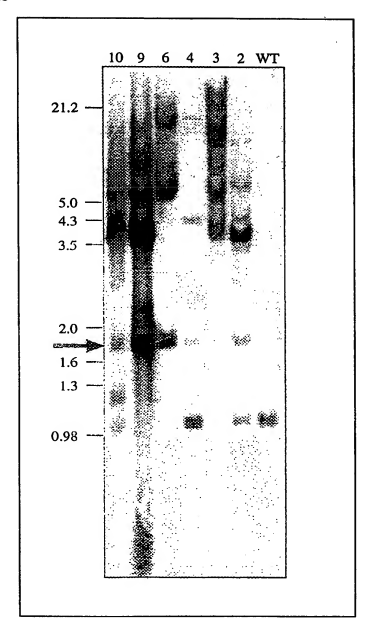
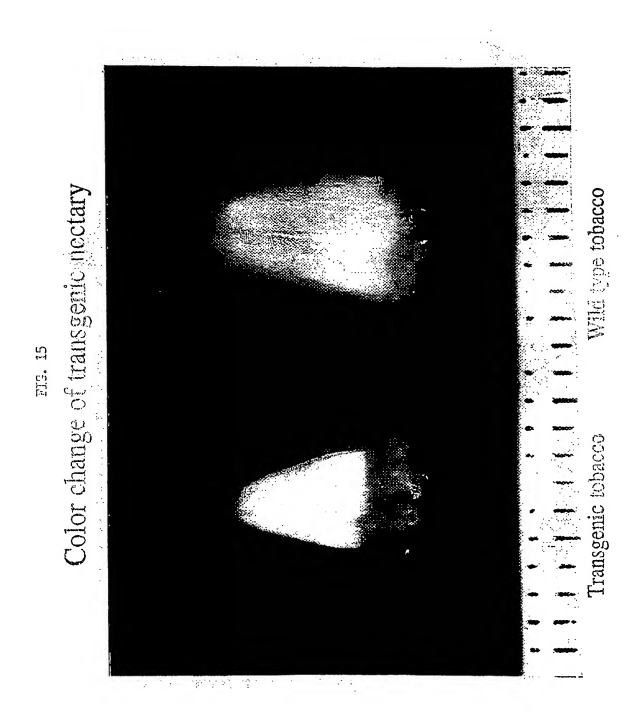


Fig. 14



SUBSTITUTE SHEET (RULE 26)

International application No. PCT/US97/17819

A. CLASSIFICATION OF SUBJECT MATTER IPC(6): C12N 9/02, 1/20, 15/09, 15/63; C12P 13/00							
115 C1 435/132 189 252.3 252.33, 419, 320.1; 536/23.2; 800/205							
According to International Patent Classification (IPC) or to both	national classification and IPC						
B. FIELDS SEARCHED	1						
Minimum documentation searched (classification system followed							
U.S. : 435/132, 189, 252.3, 252.33, 419, 320.1; 536/23.2;							
Documentation searched other than minimum documentation to the	extent that such documents are included	in the fields searched					
Electronic data base consulted during the international search (na Please See Extra Sheet.	me of data base and, where practicable	, search terms used)					
C. DOCUMENTS CONSIDERED TO BE RELEVANT							
Category* Citation of document, with indication, where ap	ppropriate, of the relevant passages	Relevant to claim No.					
cDNA for astaxanthin biosynthesis from	KAJIWARA et al. Isolation and functional identification of a novel cDNA for astaxanthin biosynthesis from Haematococcus pluvialis,						
Y and astaxanthin in Escherichia coll October 1995, Vol. 29, pages 343-352	and astaxanthin in Escherichia coli. Plant Molecular Biology. October 1995, Vol. 29, pages 343-352, see abstract.						
X LOTAN et al. Cloning and expression encoding β-C-4-oxygenase, that of ketocarotenoid canthaxanthin in Hae	convert β -carotene to the	10, 17, and 19-23					
Letters. 08 May 1995, Vol. 364, document.	pages 125-128, see entire	24-33					
X Further documents are listed in the continuation of Box							
 Special categories of cited documents: A* document defining the general state of the art which is not considered 	"T" later document published after the is date and not in conflict with the ap the principle or theory underlying	plication but cited to understand					
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L document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other	when the document is taken alone	the claimed invention cannot be					
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p document published prior to the international filing date but later than the priority date claimed							
Date of the actual completion of the international search	Date of mailing of the international	search report					
11 DECEMBER 1997	2 6 JAN 1998						
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231	Authorized officer NASHAAT T. NASHED	OPL					
Facsimile No. (703) 305-3230	Telephone No. (703) 308-0196	V					

International application No. PCT/US97/17819

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No
x Y	WO 96/06172 A1 (KIRIN BEER KABUSHIKI KAISHA) 29 February 1996, page 1, abstract.	17 2, 10, and 19-33
X, P Y, P	HARKER et al. Biosynthesis of ketocarotenoids in transgenic cyanobacteria expressing the algal gene for β-C-4-oxygenase, crtO. FEBS Letters. 03 March 1997, Vol. 404, pages 129-134, see abstract.	2, 6, 8, 21-25
A	AUSICH, R. L. Production of carotenoids by recombinant DNA technology. Pure & Appl. Chem. 1994, Vol. 66, pages 1057-1062, abstract.	24-25
A	MISAWA et al. Canthaxanthin biosynthesis by the conversion of methylene to keto groups in a hydrocarbon β-carotene by a single gene. Biochem. Biophys. Res. Comm. 26 April 1995, Vol. 209, No. 3, pages 867-876.	2, 6, 8, 10, 14, 16, and 19-33
A	DEMMIG-ADAMS, B. Carotenoids and photoprotection in plants: a role for the xanthophyll zeaxanthin. Biochem. Biophys. Acta. 1990, Vol. 1020, pages 1-24.	24-33
A	KRINSKY, N. I. Antioxidant function of carotenoids. Free Rad. Biol. Med. 1989, Vol. 7, pages 617-635.	26 and 27
A	SANDMANN, G. Carotenoid biosynthesis in microorganisms and plants. Eur. J. Biochem. 1994, Vol. 223, pages 7-24.	24 and 25

International application No. PCT/US97/17819

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2. X Claims Nos.: 1, 3-5, 7, 9, 11-13, 15 and 18 because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
The claims are drawn to specific nucleic acid and amino acid sequences of SEQ ID NO: 2 and SEQ ID NO: 4, respectively. Applicants have not provided the sequences in computer readable form to search the sequences in commercial data bases. Thus, the claims could not be meaningfully searched.
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
Please See Extra Sheet.
1. X As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
As only some of the required additional search fees were timely paid by the applicant, this international search report covers
only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
restricted to the invention first mentioned in the cannis, is a covered by cannot be a
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.

International application No. PCT/US97/17819

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, STN: Medline, Caplus, Scisearch, Lifesci, Biosis, Embase, Wpids, Agricola, and Biotechds. Search terms: haematococcus, pluvialis, DNA, cDNA, Sequence, Astaxanthin, canthaxanthin, echinenone, isocryptoxanthin, crytoxanthin, hydroxyechinenone, zeaxanthin, adonirubin, adoniranthin and crtO.

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group I, claims 2, 6, 8, 10, 14, 16, 17, and 19-27, drawn to DNA coding for an oxido-reductase which is involve in the biosynthesis of astaxanthin, the enzyme of SEQ ID NO: 4, vector, host cell and a method of making xanthophylls.

Group II, claims 28-33, drawn to a transgenic plants, and a special vector for transforming plants.

The inventions listed as Groups I and II do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: The special technical feature of the invention of Group I is the DNA coding for the oxido-reductase of SEQ ID NO: 4. Group I encompass the DNA, the enzyme, vector containing the said DNA, host cell and a single use for the DNA which is the method of producing xanthophylls. The invention of Group II is drawn to a transgenic plant containing the DNA of Group I and vectors useful for transforming plants. Thus, the invention of Group II represent a second use for the DNA. Thus, the inventions of Groups I and II are not so linked with a special technical feature as to form a single inventive concept under PCT Rule 13.1, see 37 C.F.R. 1.475(d).

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